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WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

A61K 39/00, 39/44, 39/02, 39/085, C12N
15/63, 1/20, 1/21, 15/00, C12P 21/00,
G01N 33/53, C07K 14/00

(11) International Publication Number:

WO 00/09154

(43) International Publication Date:

24 February 2000 (24.02.00)

(21) International Application Number:

PCT/US98/16766

A1

(22) International Filing Date:

13 August 1998 (13.08.98)

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Published

With international search report.

(54) Title: BACTERIAL SUPERANTIGEN VACCINES

(57) Abstract

The present invention relates to genetically attenuated superantigen toxin vaccines altered such that superantigen attributes are absent, however the superantigen is effectively recognized and a n appropriate immune response is produced. The attenuated superantigen toxins are shown to protect animals against challenge with wild type toxin. Methods of producing and using the altered superantigen toxins are described.

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Z Czech Republic LC Saint Lucia RU Komana RU Russian Federation LI Licenteratein SD Status C Dennark LK Sri Lanka SR Secretarion

TITLE OF THE INVENTION

Bacterial Superantigen Vaccines by Robert G. Ulrich Mark A. Olson

INTRODUCTION

Sina Bayari

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Staphylococcal enterotoxins (SEs) A through E are the most commom cause of food poisoning [Bergdoll, M.S.(1983) In Easom CSF, Aslam C., eds. Staphylococci and staphylococcal infections. London: Academic Press. pp 559-598] and are associated with several serious 15 diseases [Schlievert, P.M. (1993) J. Infect. Dis. 167: 997-1002; Ulrich et al. (1995) Trends Microbiol. 3: 463-468], such as bacterial arthritis [Schwab et al. (1993) J. Immunol. 150; 4151-4159; Goldenberg et 20 al. (1985) N. Engl. J. Med. 312: 764-771), other autoimmune disorders [Psnett, D. N. (1993) Semin. Immunol. 5: 65-72], and toxic shock syndrome [Schlieverst, P.M. (1986) Lancet 1: 1149-1150; Bohach et al. (1990) Crit. Rev. Microbiol. 17: 251-272]. 25 nonenterotoxic staphylococcal superantigen toxic shock syndrome toxin-1 (TSST-1) was first identified as a causative agent of menstrual-associated toxic shock syndrome [Schlievert et al. (1981) J. Infect. Dis. 143: 509-516]. Superantigen-producing Staphylococcus aureus strains are also linked to Kawasaki syndrome, 30 an inflammatory disease of children [Leung et al. (1993) Lancet 342: 1385-1388].

The staphylococcal enterotoxins A-E, toxic shock syndrome toxin-1 (TSST-1), and streptococcal pyrogenic

exotoxins A-C are soluble 23-29-kD proteins commonly referred to as bacterial superantigens (SAgs).

Bacterial superantigens are ligands for both major histocompatibility complex (MHC) class II molecules, expressed on antigen-presenting cells, and the variable portion of the T cell antigen receptor β chain (TCR Vβ) [Choi et al. (1989) Proc. Natl. Acad. Sci. USA 86:8941-8945; Fraser, J.D. (1989) Nature 339:221-223; Marrack et al. (1990) Science 248: 705-711; Herman et al. (1991) Annu. Rev. Immunol. 9: 745-772; Mollick et al. (1989) Science 244:817-820].

Each bacterial superantigen has a distinct affinity to a set of TCRVB, and coligation of the MHC class II molecule polyclonally stimulates T cells 15 [White et al. (1989) Cell 56: 27-35; Kappler et al. (1989) Science 244: 811-813; Takimoto et al. (1990) Eur J. Immunol. 140: 617-621]. Pathologically elevated levels of cytokines that are produced by activated T cells are the probable cause of toxic shock symptoms [Calson et al. (1985) Cell. Immunol. 20 96: 175-183; Stiles et al. (1993) Infect. Immun. 61: 5333-5338]. In addition, susceptibility to lethal gram-negative endotoxin shock is enhanced by several bacterial superantigens [Stiles, et al., supra]. 25 Although antibodies reactive with superantigens are present at low levels in human sera [Takei et al. (1993) J. Clin. Invest. 91: 602-607], boosting antibody titers by specific immunization may be efficacious for patients at risk for toxic shock syndrome and the other disorders of common etiology. 30

A vaccine approach to controlling bacterial superantigen-associated diseases presents a unique set of challenges. Acute exposure to bacterial superantigens produces T cell anergy, a state of

specific non-responsiveness [Kawabe et al. (1991) Nature 349: 245-248], yet T cell help is presumably a requirement for mounting an antibody response.

Presently, the only superantigen vaccines available are chemically inactivated toxoids from 5 different bacteria which have several disadvantages. The chemical inactivation process can be variable for each production lot making the product difficult to characterize. The chemical used for inactivation, (e.g. formaldehyde), is often toxic and does not negate the possibility of reversion of the inactivated superantigen to an active form. In addition, the yields of wild-type toxin from bacterial strains used for making toxoids are often low.

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SUMMARY OF THE INVENTION

The present invention relates to a vaccine which overcomes the disadvantages of the chemically inactivated toxoids described above. The superantigen 20 vaccine(s) of the present invention is/are designed to protect individuals against the pathologies resulting from exposure to one or several related staphylococcal and streptococcal toxins. The superantigen vaccine is comprised of a purified protein product that is 25 genetically attenuated by DNA methodologies such that superantigen attributes are absent, however the superantigen is effectively recognized by the immune system and an appropriate antibody response is produced.

Specifically, the vaccine of the present invention is a product of site-directed mutagenesis of the DNA coding sequences of superantigen toxins resulting in a disruption of binding to both the MHC class II receptor and to the T-cell antigen receptor. 35 A comprehensive study of the relationships of the

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superantigen structures of TSST-1, streptococcal
pyrogenic exotoxin-A (SPEa), staphylococcal
enterotoxin B (SEB), and staphylococcal enterotoxin A,
to receptor binding were undertaken to provide insight
into the design of the vaccine. From these studies,
critical amino acid residues of the toxin responsible
for binding the superantigen to the human MHC receptor
were defined. Site-directed mutagenesis of the gene
encoding the toxin and expression of the new protein
product resulted in a superantigen toxin with
disrupted binding to the MHC receptors.

Therefore, it is an object of the present invention to provide a superantigen toxin DNA fragment which has been genetically altered such that binding of the encoded altered toxin to the MHC class II or T-cell antigen receptor is disrupted. Such a DNA fragment is useful in the production of a vaccine against superantigen toxin infections.

It is another object of the present invention to provide a superantigen toxin amino acid sequence which has been altered such that the binding of the encoded altered toxin to the MHC class II or T-cell antigen receptor is disrupted. Such a sequence is useful for the production of a superantigen toxin vaccine.

It is another object of the invention to provide a recombinant vector comprising a vector and the DNA fragment described above.

It is a further object of the present invention to provide host cells transformed with the abovedescribed recombinant DNA constructs. Host cells include cells of other prokaryotic species or eukaryotic plant or animal species, including yeasts,

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fungi, plant culture, mammalian and nonmammalian cell lines, insect cells and transgenic plants or animals.

It is another object of the present invention to provide a method for producing altered superantigen 5 toxin with disrupted MHC class II and T-cell antigen receptor binding which comprises culturing a host cell under conditions such that a recombinant vector comprising a vector and the DNA fragment described above is expressed and altered superantigen toxin is thereby produced, and isolating superantigen toxin for use as a vaccine against superantigen toxin-associated bacterial infections and as a diagnostic reagent.

It is still another object of the invention to provide a purified altered superantigen toxin useful as a vaccine and as a diagnostic agent.

It is another object of the invention to provide a purified altered superantigen toxin for the therapeutic stimulation of, or other in vivo manipulation of, selective T cell subsets, or ex vivo 20 manipulation of T cells for in vivo therapeutic purposes in mammals. Diseases, such as autoimmunity, wherein T-cell responses of limited diversity (oligoclonal) are evident. Altered superantigens may be used to therapeutically inactivate (induce anergy in) T cells in diseases wherein oligoclonal T-cell 25 responses are evident such as autoimmune diseases, for example. For diseases in which specific T-cell subsets are not active or are anergetic, altered superantigens may be used to therapeutically stimulate these T cells. Such disease include, but are not limited to, infectious diseases and cancers wherein specific subsets of cytotoxic or helper T cells are inactivated or are otherwise unable to respond normally to the antigenic stimulation of the disease 35 moiety.

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It is a further object of the present invention to provide an antibody to the above-described altered superantigen toxin for use as a therapeutic agent and as, a diagnostic agent.

It is yet another object of the invention to provide a superantigen toxin vaccine comprising an altered superantigen toxin effective for the production of antigenic and immunogenic response resulting in the protection of an animal against superantigen toxin infection.

It is a further object of the invention to provide a multivalent superantigen toxin vaccine comprising altered toxins from a variety of streptococcal and staphylococcal toxins effective for the production of antigenic and immunogenic response resulting in the protection of an animal against infection with bacterial superantigen toxin-expressing strains and against other direct or indirect exposures to bacterial superantigen toxins such as might occur by ingestion, inhalation, injection, transdermal or other means.

It is yet another object of the present invention to provide a method for the diagnosis of superantigen toxin-associated bacterial infection comprising the steps of:

- (i) contacting a sample from an individual suspected of having a superantigen toxin-associated bacterial infection with antibodies which recognize superantigen toxin using antibodies generated from the altered superantigen toxin; and
- (ii) detecting the presence or absence of a superantigen-associated bacterial infection by detecting the presence or absence of a complex formed between superantigen toxin in said sample and antibodies specific therefor.

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It is yet another object of the present invention to provide a method for the diagnosis of superantigen bacterial infection comprising the steps of:

- (i) contacting a sample from an individual suspected of having the disease with lymphocytes which recognize superantigen toxin produced by said superantigen bacteria or lymphocytes which recognize altered superantigen toxin; and
- (ii) detecting the presence or absence of responses of lymphocytes resulting from recognition of 10 superantigen toxin. Responses can be, for example, measured cytokine release, increase of activation markers, mitotic activity, or cell lysis. lymphocytes responding to the altered superantigen 15 toxins recognize them as recall antigens not as superantigens, therefore the response is an indicator of prior exposure to the specific superantigen. absence of a response may indicate no prior exposure, a defective immune response or in some cases a manifestation of T-cell anergy. Anergy is defined 20 here as antigen-specific or a generalized nonresponsiveness of subsets of T cells.

It is a further object of the present invention to provide a diagnostic kit comprising an antibody against an altered superantigen toxin and ancillary reagents suitable for use in detecting the presence of superantigen toxin in animal tissue or serum.

It is another object of the present invention to provide a detection method for detecting superantigen toxins or antibodies to superantigen toxin in samples, said method comprising employing a biosensor approach. Such methods are known in the art and are described for example in Karlsson et al. (1991) J. Immunol. Methods 145, 229-240 and Jonsson et al. (1991) Biotechniques 11, 620-627.

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It is yet another object of the present invention to provide a therapeutic method for the treatment or amelioration of symptoms of superantigen-associated bacterial infection, said method comprising providing to an individual in need of such treatment an effective amount of sera from individuals immunized with one or more altered superantigen toxins from different bacteria in a pharmaceutically acceptable excipient.

10 It is further another object of the present invention to provide a therapeutic method for the treatment or amelioration of symptoms of superantigen toxin-associated bacterial infection, said method comprising providing to an individual in need of such 15 treatment an effective amount of antibodies against altered superantigen toxins in a pharmaceutically acceptable excipient.

It is another object of the present invention to provide a therapeutic method for the treatment or amelioration of symptoms of bacterial superantigen toxin infection, said method comprising providing to an individual in need of such treatment an effective amount of altered superantigen from a variety of streptococcal and staphylococcal bacteria in order to inhibit adhesion of superantigen bacterial toxin to MHC class II or T-cell receptors by competitive inhibition of these interactions.

It is yet another object of the present invention to provide a therapeutic method for the treatment of diseases that may not be associated directly with superantigen toxins but which result in specific nonresponsiveness of T-cell subsets, said method comprising the administration of altered superantigen toxins, in vivo or ex vivo, such that T-cell subsets are expanded or stimulated. Diseases which cause

anergy or nonresponsiveness of T-cells include, but are not limited to, infectious diseases.

It is another object of the present invention to provide a therapeutic method for the treatment of diseases associated with expanded or over-stimulated T-cell subsets, such as autoimmunity for example, said method comprising administration of altered superantigen toxin, in vivo or ex vivo, such that anergy or inactivation of disease associated T-cells is produced. In this case, superantigen mutants can be designed with altered but not attenuated T-cell receptor binding, to cause anergy of only the select (i.e. 1-3) T-cell subsets that are pathologically activated.

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BRIEF DESCRIPTION OF THE DRAWINGS

These and other features, aspects, and advantages of the present invention will become better understood with reference to the following description and appended claims, and accompanying drawings where:

Figure 1. Staphylococcal and streptococcal superantigen amino acid sequence homologies, compiled with Genetics Computers Group of Univ. of Wisconsin software.

Figure 2. Comparison of mutant SEB and SEA biological activities.

A. SEB mutant HLA-DR1-binding; B. SEA mutant HLA-DR1-binding; C. T-cell recognition of SEA and SEB mutants. Binding of bacterial superantigens to cell surface DR1 was measured by laser fluorescence-activated flow cytometry. A representative experiment of three performed is shown. The mutants SEA D197N, the homologous SEB D199N, and SEA L11Y had no effect on binding or T-cell recognition, and were used for

controls. Human T-cell proliferation, assessed by [³H]thymidine incorporation, was measured in response to SEA (Y64A) or SEB (Y61A) mutants and controls that retained DR1-binding affinities. Each data point represents the mean of triplicate determinations; SEM <5%.

Figure 3. Sequence and secondary structural alignment of bacterial: superantigen toxins. Analyses were performed with the applications PILEUP and PROFILE from the Computer Genetics Group (Madison, WI) using sequence data obtained from a variety of sources. Amino acid residue numbering is based on the SEA sequence.

Figure 4. Detection of TNF- α (a), IL-1 α (B), IL-15 6 (C) and IFN- γ (D) in the serum of mice injected with SEA (open circles), LPS (open triangles), or SEA plus LPS (open squares). Values for TNF- α and IL-1 α represent the mean of duplicate samples, with an SEM of \pm 5%. INF- γ and IL-6 values represent the mean of duplicate and triplicate samples, respectively. The SEMs for IFN- γ and IL-6 readings were \pm 5% and \pm 10%, respectively.

Figure 5. Mutant SEA vaccines that have attenuated major histocompatibility complex class II

25 or T-cell antigen receptor binding do not induce T-cell anergy. Mice were given three doses of wild type (WT) SEA or site-specific mutant vaccine, plus adjuvant. Control animals received adjuvant alone or were untreated; 2 weeks after final injection, pooled

30 mononuclear cells were collected from spleens of 4 mice from each group. Results are represented as mean cpm (±SD) of quadruplicate wells incubated with 100 ng/ml WT SEA for 72 h and then pulse-labeled for 12 h with [3H]thymidine. P<0.0001 (analysis of variance

for repeated measures comparing untreated, adjuvant, Y64A, and Y92A to WT SEA group).

Figure 6. No superantigen-induced T-cell anergy is exhibited by rhesus monkeys immunized with the vaccine B899445. Peripheral blood lymphocytes were incubated with titrated concentrations of wild-type superantigens from individual rhesus monkeys (K422 and N103) that were immunized with B899445. T-cell proliferation was assessed by [3H]thymidine incorporation. Each data point represents the mean of triplicate determinations; SEM <5%.

Figure 7. Antibody responses of rhesus monkeys immunized with a combined vaccine consisting of B899445 (SEB) and A489270 (SEA). The antibody levels were measured by ELISA, using plates coated with SEA, SEB or SEC1 as listed. Monkey G8 is a non-immunized control. SEM <5% for triplicate measurements.

DETAILED DESCRIPTION

20 The present invention relates in part to a vaccine against superantigen toxin-associated bacterial diseases. The superantigen vaccines used in this study were developed by engineering changes in the receptor-binding portions of superantigen toxins 25 to reduce receptor-binding affinities and toxicity while maintaining antigenicity.

Five different superantigen vaccines are described in this application: staphylococcal enterotoxin A, staphylococcal enterotoxin B,

30 staphylococcal enterotoxin C1, toxic-shock syndrome toxin-1, and streptococcal pyrogenic exotoxin-A. For each of the superantigen toxins above, a comprehensive study of the relationships of the toxin protein structure to receptor binding was undertaken to

35 provide insight into the design of the vaccine. The

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study employed site-directed mutagenesis of toxin and receptor molecules, molecular modeling, protein structure and binding studies. Following these studies, toxins were altered by site-directed mutagenesis to attenuate MHC class II binding and biological activity to an essentially non-specific level. The engineered vaccines were evaluated at each stage of analysis to determine mouse and human T-cell reactivities in vitro, serological responses and toxicity in mice and monkeys.

In one embodiment, the present invention relates to an altered superantigen toxin protein having an amino acid sequence which has been altered such that the binding of the toxin to the MHC class II receptor is disrupted.

Comparison of amino acid sequences (Fig. 1) suggested that bacterial superantigens fall into groups consisting of (1) SEA, SED and SEE, (2) SEB, staphylococcal enterotoxins C1-C3 (SEC1-3), the 20 streptococcal pyrogenic exotoxins A (SPE-A) and C (SPE-C), (3) TSST-1 and (4) the exfoliative toxins (ETA, ETB) and streptococcal pyrogenic exotoxin B (SPE-B), which are the most distant from the others in sequence. Although not available to the inventor when 25 the inventions were first conceived and proof of principle was obtained, the x-ray crystallographic structures of several bacterial superantigens are now Diverse superantigens, such as SEB and TSST-1, known. appear to have little sequence in common, yet they 30 exhibit homologous protein folds composed largely of β strands [Prasad, G.S. et al. (1993) Biochemistry 32, 13761-13766; Acharya, R.K. et al. (1994) Nature 367, 94-97; Swaminathan, S. et al. (1992) Nature 359, 801-806] within two distinct domains. Differences between the proteins are located primarily in highly variable 35

regions comprised of several surface loops, such as the disulfide-bonded loop which is absent from TSST-1 and at the amino terminus.

The X-ray crystal structures of SEB and TSST-1 complexed with HLA DR1 are known [Kim, J. et al. (1994) Science 266, 1870-1874; Jardetzky, T.S. et al. (1994) Nature 368, 711-718]. The region of HLA DR1 that contacts SEB consists exclusively of α subunit surfaces. The main regions of SEB involved are two conserved sites: a polar pocket derived from three β 10 strands of the \$\beta\$ barrel domain and a highly solventexposed hydrophobic reverse turn. The polar binding pocket of SEB contains a glutamate and two tyrosines that accommodate Lys39 of the a subunit of HLA DR1, while the hydrophobic region consists of a leucine and flanking residues that make several contacts with the HLA DRa chain. The HLA DR1 binding sites of both TSST-1 and SEB overlap significantly. The hydrophobic binding contacts of other SAg with the HLA $DR\alpha$ chain 20 have been proposed [Ulrich, et al. (1995). Nature, Struct. Biol 2, 554-560] to be similar to those found in SEB and TSST-1. A motif consisting of a leucine in a reverse turn [Ulrich et al. (1995), supra] is conserved among bacterial superantigens and may provide the key determinant (hydrophobic or otherwise) for binding HLA-DR. However, TSST-1 does not have a highly charged residue in the polar pocket that interacts with Lys39 of the HLA DRα chain and uses an alternative conformational binding mode that allows TSST-1 to interact with HLA DR1 β -chain residues and 30 the carboxy-terminal region of the antigenic peptide.

Both SEA and SEE bind to the β subunit of DR by means of a single zinc atom [Fraser, J.D. et al. (1992) Proc. Natl. Acad. Sci. USA 89, 5507-5511]. The amino-terminal domain of SEA interfaces with the HLA

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DRα chain [Ulrich, et al. (1995)], while SEA C-terminal domain residues His187, His225 and Asp227 form a zinc-coordination complex, likely with His-81 from the β chain of an adjoining HLA DR molecule. However, our results have shown that binding of superantigen to the HLA DRβ subunit does not directly stimulate T cells [Ulrich et al. (1995) Nature, Struct. Biol. 2, 554-560], but increases the potential of the bound SEA to interact with the α chain of another HLA DR, thus increasing the biological potency.

A least-squares superimposition of the unbound molecules of modeled SEA and the crystal structure of SEB, aligned according to their structurally conserved 15 α-helical and β-strand regions, exhibited a global folding pattern which is very similar. Differences between the two structures are calculated to be located primarily in loops of low sequence homologies, with the largest positional deviations occurring 20 between structurally conserved regions of residues 18-20, 30-32, 173-181, 191-194, and the cysteine-loop region (90-111). Only one of these regions in SEB makes significant contact (residue Y94 [Y=tyrosine] in particular) with the HLA-DR1 molecule [Jardetzky, T.S. et al. (1994) Nature 368, 711-718].

The binding interface between SEB and HLA-DR1 consists principally of two structurally conserved surfaces located in the N-terminal domain: a polar binding pocket derived from three ß-strand elements of the ß-barrel domain and a hydrophobic reverse turn. The binding pocket of SEB contains residues E67 (E=Glutamic acid), Y89 (Y=Tyrosine) and Y115 (Y=tyrosine), and binds K39 (K=Lysine) of the DRa subunit. The amino acid one letter code is defined as the following: A= Alanine (Ala), I= Isoleucine (Ile),

WO 00/09154 PCT/US98/16766

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L= Leucine (Leu), M= Methionine (Met), F= Phenylalanine (Phe), P= Proline (Pro), W=Tryptophan (Trp), V=Valine (Val), N= Asparagine (Asn), C=Cysteine (Cvs), O= Glutamine (O), G= Glycine (Gly), S= Serine 5 (Ser), T= Threonine (Thr), Y= Tyrosine (Tyr), R= Arginine (Arg), H=Histidine (His), K= Lysine (Lys), D= Aspartic acid (Asp), and E= Glutamic acid (Glu). For SEA, the binding interface with the DR molecule is modeled to contain a similar binding pocket consisting of residues D70, Y92 and Y108. Mutation of residue 10 y89 in SEB or Y92 in SEA to alanine (Fig. 2) resulted in greater than 100-fold reduction in DR1 binding. The substitution of alanine for Y89 in SEB and Y92 in SEA eliminates the hydrogen bond with K39 and disrupts packing interactions with adjacent protein residues. 15 Modeling of the SEA mutant Y92A predicts an increase in solvent-accessible surface area for Y108 by a factor of two greater than the wild-type structure, allowing the formation of a hydrogen bond to the carboxylate group of D70 and thus disrupting key 20 anchoring and recognition points for HLA-DR1. This effect is expected to be somewhat less in SEB due to the longer side chain at E67. Substitution of SEB Y115 with alanine also resulted in greater than 100fold reduction of binding. In contrast, the same 25 replacement of Y108 in SEA yielded little to no change in DR1 binding (Fig. 2a), suggesting the primary importance of SEA residues Y92 and D70 for stabilizing interactions with K39. The K39 side chain of DRa 30 forms a strong ion-pair interaction with the SEB E67 carboxylate group and hydrogen bonds with the hydroxyl groups of Y89 and Y115. Substitution of SEB E67 by glutamine reduced binding affinity by greater than 100-fold (Fig. 2), reflecting the replacement of the strong ionic bond with a weaker hydrogen bond.

optimize ion-pair interactions of the analogous SEA site, the shorter carboxylate side chain of D70 is predicted to shift K39 of DRα, weakening interactions with SEA Y108. The substitution of alanine for SEA Y108 is thus more easily accommodated than the homologous substitution of SEB Y115, without loss in DR1 binding.

Comparisons of the polar pocket with other bacterial superantigens were then made. SEC1-3 and 10 SPE-A have conserved the critical DR1 bindinginterface residues (Fig. 1), and share with SEB and SEA secondary structural elements of the DR1-binding surfaces. Asparagine in SED (N70) replaces the acidic side chain present in SEA, SEB, SPE-A and SEC1-3. Accordingly, for SED the salt bridge of the polar pocket is likely to be replaced by a hydrogen bond. Overall, DR1 affinities for SED and SEA appeared to be equivalent (Fig. 2b), indicating that other interactions may compensate for the absence in SED of the ion-pair found in the other superantigens. For 20 the case of TSST-1, mutating DR α residues K39 to serine or M36 to isoleucine has been shown to greatly reduce binding [Panina-Bordignon et al. (1992) J. Exp. Med. 176: 1779-1784]. Although primarily 25 hydrophobic, the critical TSST-1 structural elements are conserved with the SEA and SEB polar binding pocket. SEB residues Y89 and Y115 are homologous to T69 and I85 in TSST-1, respectively, and SEB E67 is replaced by I46. These TSST-1 residues are positioned 30 in a conserved G-barrel domain found in both SEB and However, the TSST-1 site lacks polarity equivalent to SEB/SEA, and hydrogen bonding with the hydroxyl of TSST-1 residue T69 would require that DRa K39 extend 5 Å into the pocket. TSST-1 binding utilizes an alternative strategy [Kim et al. (1994) 35

Science 266:1870-1874] consisting of hydrophobic contacts centered around residue I46, and potential ionic or hydrogen bonds bridging DRα residues E71 and K67 to R34 and D27, respectively, of TSST-1.

5 The hydrophobic region of the binding interface between SEB and the HLA-DR1 molecule consists of SEB residues 44-47, located in a large reverse turn connecting &-strands 1 and 2 of SEB. These residues appear to make strong electrostatic interactions with 10 DRa through their backbone atoms. The mutation of L45 to an arginine reduced overall HLA-DR1 binding greater than 100-fold (Fig. 2b), attributable to the less energetically favorable insertion of a highly charged residue into a hydrophobic depression on the DR1 15 molecule. The modeled DR1-SEA complex presents similar interactions with the SEA backbone atoms, with the exception of a glutamine (Q49) replacing SEB Y46. Mutation of L48 to glycine in SEA (homologous to L45 of SEB) has been reported to decrease T-cell 20 responses. SEB L45 and the comparable L30 of TSST-1 are the most extensively buried residues in the DR1 interface. The leucine is conserved among the bacterial superantigens (Fig. 3) and may provide the necessary hydrophobic structural element for surface 25 complementarity with DR1, consistent with the mutagenesis data for SEB and SEA.

The inventor has performed similar structure and function studies with TSST-1. SEC1 and SPE-A.

In determining the overall affinity of the

30 superantigen for DR1, a contributory role is played by
structural variations around the common binding
motifs. A short, variable structured, disulfidebonded loop is found in SEA and a homologous longer
loop in SEB. The SEB residue Y94, contained within

35 this loop, forms hydrophobic interactions with L60 and

A61 of the DRa subunit. Replacement of Y94 with alanine partially inhibits DR1 binding (Fig. 2a.b). An alanine is found in SEA (A97) and SEE at the position equivalent to SEB Y94, and mutating this residue in SEA to tyrosine results in disrupted instead of stabilized interactions with DR1 (Fig. 2a). Although the disulfide loops differ in structure between SEA and SEB, A97 apparently contributes to the DRa binding interface in a manner similar to Y94 of 10 SEB. Because TSST-1 lacks a disulfide loop, similar contacts with DRα are replaced by interactions with βstrands of TSST-1. In a like manner, the absence of a salt bridge between the residues K39 of DRg and N65 of SED is apparently compensated for by stabilizing 15 interactions occurring outside of the otherwise conserved dominant binding surfaces (Fig. 2a).

The amino acid residues in contact with TCR are

located in regions of high sequence variability, presenting a unique surface for interaction with the 20 TCR. Residues implicated in TCR interactions by mutagenesis of SEA and SEB reside in variable loop regions, while TSST-1 mutants that affect TCR binding are mainly located in an α helix [Acharya, R.K. et al. (1994) Nature 367, 94-97; Kim, J. et al. (1994) Science 266, 1870-1874]. Specifically, mutations that 25 diminish T-cell receptor recognition of SEB include residues N23, Y61, and the homologous SEA N25 or Y64 (Fig. 2c). SEA residues S206 and N207 also control T-cell responses [Hudson, et al. (1992) J. Exp. Med. 30 177: 175-184]. Mutants of the polar binding pocket, SEA Y92A and SEB Y89A, equivalently reduced T-cell responses (Fig. 2c), reflecting the observed decreases in DR1-binding (Fig. 2a, b). While supporting reduced T-cell responses, mutants SEA Y64A

and SEB Y61A retained normal affinities for DR1 (Fig. 2a-c).

In view of the detailed description of the present invention and the results of molecular modelling and structural studies of staphylococcal and streptococcal superantigen toxins discussed above, any amino acid sequence derived from a superantigen toxin can be altered. Sequences of several superantigen toxins are already known and available to the public in sequence databases such as GenBank, for example. 10 The superantigen toxin sequence is preferably altered at the hydrophobic loop or polar binding pocket depending on the superantigen. Alternatively, residues adjacent to the hydrophobic loop or polar binding pocket that contact HLA-DR or residues at sites that can indirectly alter the structure of the hydrophobic loop or polar pocket can be altered. The number of residues which can be altered can vary, preferably the number can be 1-2, more preferably 2-3, and most preferably 3-4, or more with the limitation 20 being the ability to analyze by computational methods the consequences of introducing such mutations. residues which can be altered can be within 5 amino acid residues of the central Leucine of the 25 hydrophobic loop (such as L45 of SEB), or within 5 residues of one of the amino acid residues of the polar binding pocket that can contact HLA-DR, (such as E67, Y89, or Y115 of SEB), more preferably, within 3 amino acid residues of the central Leucine of the hydrophobic loop (such as L45 of SEB), or within 3 30 residues of one of the amino acid residues of the polar pocket that can contact HLA-DR, (such as E67, Y89, or Y115 of SEB), and most preferably, the central Leucine of the hydrophobic loop (such as L45 of SEB), 35 or one of the amino acid residues of the polar binding

pocket that can contact HLA-DR, (such as E67, Y89, or Y115 of SEB). The residues can be changed or substituted to alanine for minimal disruption of protein structure, more preferably to a residue of opposite chemical characterisitcs, such as hydrophobic to, hydrophilic, acidic to neutral amide, most preferably by introduction of a residue with a large hydrated side chain such as Arginine or Lysine. addition, side chains of certain nonconserved 10 receptor-binding surfaces, can also be altered when designing superantigen toxins with low binding affinities. These residues can include Y94 of SEB and structurally equivalent residues of other superantigens, such as A97 of SEA, or any side chain 15 within 5 residues from these positions or any side chain in discontinuous positions (discontinuous positions are defined as amino acid residues that fold together to form part of a discrete three-dimensional structural unit but are not present on the same 20 secondary structural unit e.g. α helix or β -strand) such as disulfide-bonded side chains, that involve. directly or indirectly, the nonconserved receptor contact surfaces outside of the polar binding pocket or hydrophobic loop. Further, amino acid residues 25 involved with protein folding or packing can be altered when designing superantigen toxins with low binding affinities [Sundstrom et al. (1996) EMBO J. 15, 6832-6840; Sundstrom et al. (1996) J. Biol. Chem. 271, 32212-32216; Acharya et al. (1994) Nature 367, 94-97; Prasad et al. (1993) Biochem. 32, 13761-13766; 30 Swaminathan et al. (1992) Nature 359, 801-806]. Furthermore, especially for superantigens with higher affinities for T-cell antigen receptors, side chains of amino acids within 5 residues of the position 35 represented by N23 (conserved residue in most

superantigens) , N60 (conserved Asn or Trp in most superantigens) Y91 (semiconserved hydrophobic residues Trp, Ile, Val, His in most superantigens) and D210 of SEB (conserved Asp in most superantigens) can be altered when designing superantigen toxins with low binding affinities. These residues are likely to form part of the integral molecular surfaces that are in contact with T-cell antigen receptors. Because the Tcell receptor contact areas of superantigen toxins are essential for causing specific activation or inactivation of T-cell subsets, altering residues that are unique to each superantigen but that are located within 5 residues of the positions represented by N23, N60 and Y91 can produce superantigens that affect a 15 smaller number (e.g. 1-3) of subsets. Such altered superantigen toxins can be useful as therapeutic agents.

In another embodiment, the present invention relates to a DNA or cDNA segment which encodes a superantigen toxin such as SEA, SEB, SEC-1, SPEa, and 20 TSST-1 to name a few, the sequence of which has been altered as described above to produce a toxin protein with altered binding ability to MHC Class II and/or Tcell receptors. For SEA, the following three 25 mutations were introduced into the toxin molecule: Tyrosine at amino acid position 92 changed to alanine; Aspartic acid at amino acid position 70 changed to arginine; Leucine at amino acid position 48 changed to arginine. The reduction in binding to HLA DR is additive per mutation, though one or two mutations can 30 produce a vaccine and a combination of all three mutations in one molecule produces a better vaccine. Other substitutions can also result in reduced binding.

The B899445 vaccine consists of the following three mutations simultaneously introduced into the toxin molecule: tyrosine at amino acid position 89 changed to alanine; tyrosine at amino acid position 94 changed to alanine; leucine at amino acid position 45 changed to arginine. The altered superantigen toxins can be expressed either as a full-length propolypeptide or as a polypeptide in which the leader peptide has been deleted. The full-length expressed product (SEA vaccine, A489270P; SEB vaccine B899445P, 10 B2360210P) is secreted into the periplasmic space of E. coli host cells, and the leader peptide is recognized and cleaved by a native bacterial enzymatic mechanism. The altered superantigen toxins in which 15 the leader peptide has been deleted (A489270C, B899445C), the first residue of the mature protein is encoded by the transcriptional start site and codon for methionine (ATG), and the protein is expressed as a nonsecreted product within the host E. coli cell. 20 For the TSST-1 vaccine TST30, the leucine at position 30 was changed to arginine. For the SEC1 vaccine, SEC45, the leucine at position 45 was changed to arginine. For the SPE-A vaccine, SPEA42, the leucine at position 42 was changed to arginine.

In another embodiment, the present invention relates to a recombinant DNA molecule that includes a vector and a DNA sequence as described above. The vector can take the form of a plasmid such as any broad host range expression vector for example

30 pUC18/19, pSE380, pHIL, pET21/24 and others known in the art. The DNA sequence is preferably functionally linked to a promoter such that the gene is expressed when present in an expression system and an altered superantigen toxin is produced. The expression system can be an in vitro expression system or host cells

such as prokaryotic cells, or in vivo such as DNA vaccines.

In a further embodiment, the present invention relates to host cells stably or transiently transformed or transfected with the above-described recombinant DNA constructs. The host can be any eukaryotic or prokaryotic cell including but not limited in E. coli DH5a or BL21. The vector containing the altered superantigen toxin gene is expressed in the host cell and the product of the 10 altered toxin gene, whether a secreted mature protein or a cytoplasmic product, can be used as a vaccine or as a reagent in diagnostic assays or detection methods, or for therapeutic purposes. Please see 15 e.g., Maniatis, Fitsch and Sambrook, Molecular Cloning: A Laboratory Manual (1982) or DNA Cloning, Volumes I and II (D. N. Glover ed. 1985) for general cloning methods. The DNA sequence can be present in the vector operably linked to a highly purified IgG molecule, an adjuvant, a carrier, or an agent for aid in purification of altered toxin. The transformed or transfected host cells can be used as a source of DNA sequences described above. When the recombinant molecule takes the form of an expression system, the transformed or transfected cells can be used as a 25 source of the altered toxin described above.

A recombinant or derived altered superantigen toxin is not necessarily translated from a designated nucleic acid sequence; it may be generated in any manner, including for example, chemical synthesis, or expression of a recombinant expression system. In addition the altered toxin can be fused to other proteins or polypeptides for directing transport for example into the periplasm or for secretion from the

cell. This includes fusion of the recombinant or derived altered superantigen to other vaccines or sequences designed to aid in purification, such as Hi\$-tagged, epitope-tagged or antibody Fc-fusions.

5 In a further embodiment, the present invention relates to a method of producing altered superantigen toxin which includes culturing the above-described host cells, under conditions such that the DNA fragment is expressed and a superantigen toxin protein 10 is produced. The superantigen toxin can then be isolated and purified using methodology well known in the art such as immunoaffinity chromatography or preparative isoelectric focusing. However, the method of purification is not critical to the performance of 15 the vaccine. The altered superantigen toxin can be used as a vaccine for immunity against infection with bacterial superantigen toxins or as a diagnostic tool for detection of superantigen toxin-associated disease or bacterial infection. The transformed host cells 20 can be used to analyze the effectiveness of drugs and agents which affect the binding of superantigens to MHC class II or T-cell antigen receptors. Chemically derived agents, host proteins or other proteins which result in the down-regulation or alteration of 25 expression of superantigen toxins or affect the binding affinity of superantigen toxins to their receptors can be detected and analyzed. A method for testing the effectiveness of a drug or agent capable of altering the binding of superantigen toxins to 30 their receptors can be for example computer-aided rational design or combinatorial library screening, such as phage display technology.

In another embodiment, the present invention relates to antibodies specific for the above-described altered superantigen toxins. For instance, an

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antibody can be raised against the complete toxin or against a portion thereof. Persons with ordinary skill in the art using standard methodology can raise mohoclonal and polyclonal antibodies to the altered superantigens of the present invention, or a unique portion of the altered superantigen. Materials and methods for producing antibodies are well known in the art (see for example Goding, in, Monoclonal Antibodies: Principles and Practice, Chapter 4, 1986). The antibodies can be used in diagnostic assays for 10 detection of superantigen toxin-associated infection. Neutralizing antibodies can be used in a therapeutic composition for the treatment of amelioration of anergy and/or for the treatment of a superantigen. toxin-associated infection.

In a further embodiment, the present invention relates to a method for detecting the presence of superantigen-associated bacterial infections in a sample. Using standard methodology well known in the 20 art, a diagnostic assay can be constructed by coating on a surface (i.e. a solid support) for example, a microtitration plate or a membrane (e.g. nitrocellulose membrane), all or a unique portion of the altered superantigen described above, and 25 contacting it with the serum of a person suspected of having a superantigen-associated bacterial infection. The presence of a resulting complex formed between the altered superantigen toxin and antibodies specific therefor in the serum can be detected by any of the known methods common in the art, such as fluorescent 30 antibody spectroscopy or colorimetry. This method of detection can be used, for example, for the diagnosis of superantigen-associated bacterial infections.

In yet another embodiment, the present invention relates to a method for detecting the presence of superantigen toxin in a sample. Using standard methodology well known in the art, a diagnostic assay can be constructed by coating on a surface (i.e. a solid support) for example, a microtitration plate or a membrane (e.g. nitrocellulose membrane), antibodies specific for altered superantigen toxin, and contacting it with serum or tissue sample of a person suspected of having superantigen-associated bacterial infection. The presence of a resulting complex formed between toxin in the serum and antibodies specific therefor can be detected by any of the known methods common in the art, such as fluorescent antibody 15 spectroscopy or colorimetry. This method of detection can be used, for example, for the diagnosis of superantigen-associated bacterial infection or disease such as food poisoning and toxic-shock syndrome or the detection of superantigen toxin in food and drink.

In another embodiment, the present invention relates to a diagnostic kit which contains altered superantigen toxin from a specific bacteria or several different superantigen toxins from bacteria and ancillary reagents that are well known in the art and 25 that are suitable for use in detecting the presence of antibodies to superantigen toxin-associated bacteria in serum or a tissue sample. Tissue samples contemplated can be avian, fish, or mammal including monkey and human.

30 In yet another embodiment, the present invention relates to a vaccine for protection against superantigen toxin-associated bacterial infections. The vaccine can comprise one or a mixture of individual altered superantigen toxins, or a portion thereof. When a mixture of two or more different 35

altered superantigen toxin from different bacteria is used, the vaccine is referred to as a multivalent bacterial superantigen vaccine. The vaccine is designed to protect against the pathologies resulting from exposure to one or several related staphylococcal and streptococcal toxins. In addition, the protein or polypeptide can be fused or absorbed to other proteins or polypeptides which increase its antigenicity, thereby producing higher titers of neutralizing antibody when used as a vaccine. Examples of such proteins or polypeptides include any adjuvants or carriers safe for human use, such as aluminum hydroxide.

The staphylococcal enterotoxin (SE) serotypes SEA, SED, and SEE are closely related by amino acid 15 sequence, while SEB, SEC1, SEC2, SEC3, and the streptococcal pyrogenic exotoxins B share key amino acid residues with the other toxins, but exhibit only weak sequence homology overall. However, there are considerable similarities in the known three-20 dimensional structures of SEA, SEB, SEC1, SEC3, and TSST-1. Because of this structural similarity, it is likely that polyclonal antibodies obtained from mice immunized with each SE or TSST-1 exhibit a low to high degree of cross-reaction. In the mouse, these 25 antibody cross-reactions are sufficient to neutralize the toxicity of most other SE/TSST-1, depending upon the challenge dose. For example, immunization with a mixture of SEA, SEB, TSST-1 and SPEa was sufficient to provide antibody protection from a challenge with any of the component toxins, singly or in combination.

The likelihood of substantial antigen-crossreactivity suggests that it may be possible to obtain immune protection for other (or perhaps all) staphylococcal superantigens by use of a minimal mixed composition of vaccines. For the case of staphylococcal superantigens, a combination of the component vaccines from SEA, SEB, SEC-1 and TSST-1 should be sufficient to provide immune protection

5 against SEA, SEB, SEC1-3, and TSST-1. The addition of SPEa component to the trivalent mixture will allow for sufficient protection against the streptococcal toxins SPEa and SPEc. Therefore, a multivalent vaccine consisting of the altered superantigen toxins from SEA, SEB, SEC-1, TSST-1, and SPEa as described above, is predicted to provide protective immunity against the majority of bacterial superantigen toxins.

The vaccine can be prepared by inducing expression of a recombinant expression vector 15 comprising the gene for the altered toxin described above. The purified solution is prepared for administration to mammals by methods known in the art, which can include filtering to sterilize the solution. diluting the solution, adding an adjuvant and 20 stabilizing the solution. The vaccine can be lyophilized to produce a vaccine against superantigen toxin-associated bacteria in a dried form for ease in transportation and storage. Further, the vaccine may be prepared in the form of a mixed vaccine which contains the altered superantigen toxin(s) described above and at least one other antigen as long as the added antigen does not interfere with the effectiveness of the vaccine and the side effects and adverse reactions, if any, are not increased 30 additively or synergistically. Furthermore, the vaccine may be administered by a bacterial delivery system and displayed by a recombinant host cell such as Salmonella spp, Shigella spp, Streptococcus spp. Methods for introducing recombinant vectors into host 35 cells and introducing host cells as a DNA delivery

system are known in the art [Harokopakis et al. (1997) Infect. Immun. 65, 1445-1454; Anderson et al. (1996) Vaccine 14, 1384-1390; Medaglini et al. (1995) Proc. Natl. Acad. Sci. U.S.A. 92, 6868-6872].

The vaccine may be stored in a sealed vial, ampule or the like. The present vaccine can generally be administered in the form of a liquid or suspension. In the case where the vaccine is in a dried form, the vaccine is dissolved or suspended in sterilized distilled water before administration. Generally, the vaccine may be administered orally, subcutaneously, intradermally or intramuscularly but preferably intranasally in a dose effective for the production of neutralizing antibody and protection from infection or disease.

In another embodiment, the present invention relates to a method of reducing superantigenassociated bacterial infection symptoms in a patient by administering to said patient an effective amount 20 of anti-altered superantigen toxin antibodies, as described above. When providing a patient with antisuperantigen toxin antibodies, or agents capable of inhibiting superantigen function to a recipient patient, the dosage of administered agent will vary depending upon such factors as the patient's age, 25 weight, height, sex, general medical condition. previous medical history, etc. In general, it is desirable to provide the recipient with a dosage of the above compounds which is in the range of from 30 about 1pg/kg to 10 mg/kg (body weight of patient). although a lower or higher dosage may be administered.

In a further embodiment, the present invention relates to a therapeutic method for the treatment of diseases that may not be associated directly with superantigen toxins but which result in specific nonresponsiveness of T-cell subsets or detection of abnormally low level of subsets in peripheral blood, said method comprising the administration of altered suberantigen toxins, in vivo or ex vivo, such that T-cell subsets are expanded or stimulated. Diseases which cause anergy or nonresponsiveness of T-cells include, but are not limited to, infectious diseases and cancers. The desired clinical outcome such as an increase in detectable T cell subsets or in stimulation ex vivo of T-cells through their antigen receptors, such as by antigen or anti-CD3 antibody can be measured by standard clinical immunology laboratory assays.

In yet another embodiment, the present invention relates to a therapeutic method for the treatment of 15 diseases associated with expanded or over-stimulated T-cell subsets, such as autoimmunity for example, said method comprising administration in vivo or ex vivo, of superantigen toxin altered in such a manner that only limited (1-3) T-cell subsets are stimulated but 20 that MHC class II binding affinity still remains, such that anergy or inactivation of T-cells is produced. The desired clinical outcome can be measured as a reduction of circulating blood T-cells of the targeted 25 subset(s) or diminished antigen or other antigen receptor-mediated-stimulatory responses by assays known in the art.

Described below are examples of the present invention which are provided only for illustrative purposes, and not to limit the scope of the present invention. In light of the present disclosure, numerous embodiments within the scopy of the claims will be apparent to those of ordinary skill in the art.

The following Materials and Methods were used in the Examples that follow.

Structural comparisons

Primary protein structure data are available for several bacterial superantigens, including SEA, SED, SEB, SEC1-3, TSST-1. Superantigens for which structures were unavailable were modeled using comparative techniques (HOMOLOGY program; Biosym Technologies, Inc., San Diego, CA). Before x-ray 10 crystallography data was available, SEA was modeled by using this method, and the model was in very close agreement with the experimentally determined structure. As an example, the amino acid sequence for SEA was aligned with the known structure of free and HLA-DR1 bound SEB, and the SEA molecule was built for 15 both free and DR1-bound proteins. Loop segments of SEA were generated by a de novo method. Refinement of the modeled structures was carried out by means of molecular-dynamics simulations (DISCOVER, Biosym). 20 The constructed free SEA molecule was immersed in a 5-Å layer of solvent water and the α -carbon atoms lying in the structurally conserved regions were tethered to their initial positions during the simulations. the bound SEA molecule, simulations were carried out by constructing an active-site region composed of part 25 of the SEA molecule and the DR1 molecule inside a 10-Å interface boundary, as derived from the crystal structure of the DR1-SEB complex. Amino acid residues lying in the outer boundary were rigidly restrained at their initial positions. The active-site region was immersed in a 5-A layer of water. Protein interactions were modeled by employing the consistent valence force field with a non-bonded cutoff distance of 11.0 Å. Simulations were initiated with 100 cycles 35 of minimization using a steepest descent algorithm

followed by 100-ps relaxation (using a 1.0 fs timestep). Structural comparisons between SEB, SEC1, and TSST-1 were performed by using the crystal structures (Brookhaven data holdings) aligned according to common secondary structural elements and/or by sequence and structural homology modeling.

Site-specific mutagenesis

Site-specific mutagenesis was performed according 10 to the method developed by Kunkel, using gene templates isolated from Staphylococcus aureus strains expressing SEA (FDA196E, a clinical isolate, Fraser, J.D. (1994) Nature 368: 711-718), SEB (14458, clinical isolate), SEC1 (Toxin Technologies, Sarasota, 15 FL), TSST-1 (pRN6550 cloned product, a clinical isolate, Kreiswirth, B. N. et al. (1987) Mol. Gen. Genet. 208, 84-87), and SPEa (Toxin Technologies), respectively. Modified T7 polymerase (Sequenase, U.S. Biochemical Corp., Cleveland, OH) was used to synthesize second-strand DNA from synthetic oligonucleotides harboring the altered codon and single-stranded, uracil-enriched M13 templates. Mutagenized DNA was selected by transforming E. coli strain JM101. Alternatively, double stranded DNA was used as template for mutagenesis. Mutagenized 25 sequences were confirmed by DNA sequencing (Sanger et al., 1977, Proc. Natl. Acad. Sci. USA 74: 5463-5467; Sambrook et al., 1989) using synthetic primers derived from known sequences, or universal primers. complete coding sequences were inserted into 30 expression plasmids such as pUC19, pSE380 or pET21 for production in E. coli hosts.

Protein purifications

The appropriate E. coli hosts were transformed with plasmids harboring the mutant toxin genes. In general, the bacteria were grown to an A600 0.5-0.6 in Terrific Broth (Difco Laboratories, Detroit, MI) 5 containing 50 µg/mL ampicillin or kanamycin. Recombinant proteins were induced with isopropyl-ß-Dthio-galactopyranoside (Life Technologies, Gaithersburg, MD) and recovered as cytoplasmic or bacterial periplasmic secretion products. Bacteria 10 were collected by centrifugation, washed with 30 mM NaCl, 10 mM TRIS (pH 7.6), and pelleted by centrifugation and either lysed or osmotically shocked for collection of secreted proteins. Preparations 15 were isolated by CM Sepharose ion-exchange chromatography, rabbit antibody (Toxin Technologies, Sarasota, FL) affinity columns, ion exchange HPLC or similar methods. In some cases partially purified superantigen was further purified by preparative isoelectric focusing (MinipHor; Rainin Instrument 20 Company, Inc., Woburn, MA.). The MinipHor was loaded with the SEA-enriched fraction from CM Sepharose chromatography in a solution containing 10% (v/v)glycerol and 1% (v/v) pH 6-8 ampholytes (Protein 25 Technologies, Inc., Tucson, AZ). The protein preparations were allowed to focus until equilibrium was reached (approximately 4 hr, 4°C). Twenty focused fractions were collected and aliquots of each were analyzed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting. The SEA-containing 30 fractions were pooled, and refocused for an additional 4 h. The fractions containing purified SEA were pooled and dialyzed first against 1 M NaCl (48 h, 4°C) to remove ampholytes, and then against PBS (12 h, 4°C). Legitimate amino-terminal residues were 35

confirmed by protein sequencing. Precise measurements of protein concentrations were performed by immunoassay using rabbit antibody affinity-purified with the wild-type superantigens and by the

5 bicinchoninic acid method (Pierce, Rockford, IL) using wild-type protein as standards. All protein preparations were >99% pure, as judged by SDS-PAGE and Western immunoblots. In some cases, as when used for lymphocyte assays, bacterial pyrogens were removed by passing the protein preparations over Polymyxin B affinity columns.

Binding of superantigens to HLA-DR1

The DR1 homozygous, human B-lymphoblastoid cell 15 line LG2 or L cells transfected with plasmids encoding HLA-DR1aß were used in the binding experiments. Cells were incubated 40 min (37°C) with wild-type or mutant superantigen in Hanks balanced salt solution (HBSS) containing 0.5% bovine serum albumin. The cells were washed with HBSS and then incubated with 5 μg of specific rabbit antibody (Toxin Technology, Sarasota, FL) for 1 h on ice. Unbound antibody was removed, and the cells were incubated with FITC-labelled goat antirabbit IgG (Organon Teknika Corp., Durham, N.C.) on ice for 30 min. The cells were washed and analyzed by flow cytometry (FACScan; Becton Dikinson & Co... Mountain View, CA). Controls consisted of cells incubated with affinity purified anti-toxin and the FITC labelled antibody without prior addition of 30 superantigen.

Lymphocyte proliferation

Human peripheral blood mononuclear cells were purified by Ficoll-hypaque (Sigma, St. Louis, MO)

buoyant density gradient centrifugation. Genes encoding the human MHC class II molecules DR108 (DRA and DRB1*0101 cDNA [Bavari and Ulrich (1995) Infect. Immun. 63, 423-429] were cloned into the eukaryotic expression vector pRC/RSV (Invitrogen, Carlsbad, CA), and mouse L cells were stably transfected. transfectants were selected by fluorescence-activated cell sorting (EPICS C, Coulter Corp., Hialeah, FL) using rabbit anti-DRoß antisera and FITC-goat antirabbit IgG, to produce cells that expressed a high 10 level of DRαβ21. 1 x 10⁵ cells/well of a 96-well plate were irradiated (15,000 Rad), and wild-type or mutant SE, was added. After a brief incubation period (45 min, 37°C), unbound SE was rinsed from the culture plates using warm media. The cells were cultured in 15 RPMI-1640 (USAMRIID) with 5% FBS for 72 h, and pulsedlabelled for 12 h with 1µCi [3H]-thymidine (Amersham, Arlington Heights, IL). Cells were harvested onto glass fiber filters, and [3H]-thymidine incorporation 20 into the cellular DNA was measured by a liquid scintillation counter (BetaPlate, Wallac Inc., Gaithersburg, MD). Splenic mononuclear cells or human peripheral blood mononuclear cells were obtained by buoyant density centrifugation (Histopaque; Sigma 25 Chemical Comp.) and washed three times. The cells were resuspended in medium containing 5% fetal bovine serum (FBS), and 100 μ l (4 x 10 5 cells) of the cell suspension was added to triplicate wells of 96-well flat bottom plates. The mononuclear cells were 30 cultured (37°C, 5% CO₂) with WT or mutant SEA. After 3 days the cultures were pulsed (12h) with 1 µCi/well of [3H]thymidine (Amersham, Arlington Heights, IL) and incorporated radioactivity was measured by liquid scintillation.

Gel electrophoresis and immunoblotting analysis.

The protein preparations were analyzed by SDS-PAGE (12%) and stained with Coomassie Brilliant Blue R-250 (Sigma Chemical Comp. St Louis, MO) in methanol (10% v/v) acetic acid (10% v/v). The proteins separated by SDS-PAGE (not stained) were transferred to nitrocellulose membranes (Bio-Rad Lab. Inc... Melville, NY) by electroblotting, and the membranes 10 were then blocked (12 h. 4°C) with 0.2% casein in a buffer consisting of 50 mM sodium phosphate, 140 mM sodium chloride, pH 7.4 (PBS). The membrane was then incubated (1 h, 37°C, shaking) with 2 μ g/mL of affinity-purified anti-toxin antibody (Toxin Technology, Sarasota, FL) in PBS with 0.02% casein. 15 After the membranes were thoroughly washed, peroxidase-conjugated goat anti-rabbit IgG (Cappel/Organon Teknika Corp., West Chester, PA) was added (1:5,000) and the membranes were incubated for 1 20 h (37°C) with shaking. The unbound antibody was removed by washing with PBS and bound antibody was visualized by using a Bio-Rad peroxidase development kit (Biorad, Hercules, CA). For quantitation, dilutions of wild-type preparations were immobilized on nitrocellulose membranes by using a Slot-Blot 25 apparatus (Bio-Rad). The membrane was removed from the Slot-Blot apparatus and unreacted sites were blocked (12h, 4°C) with 0.2% casein in PBS. After washing once with the PBS, the membrane was incubated 30 (1h, 37°C) with 2 μg/mL rabbit affinity purified antitoxin antibody (Toxin Technology) in PBS that contained 0.02% casein. After four washes, the bound rabbit antibody was reacted with goat anti-rabbit IgG conjugated with horseradish peroxidase (1 h, 37°C) and 35 the blots were developed using enhanced

chemiluminescence (ECL; Amersham Life Sciences,
Arlington Heights, IL) or similar methods. The amount
of mutant protein was measured by densitometry (NIH
Image 1.57 software, National Institutes of Health,

Bethesda, MD) of exposed X-ray film. Standard curves
were prepared by plotting the mean of duplicate
densitometric readings for each dilution of toxin
standard. The resulting values were fitted to a
straight line by linear regression. Concentrations of
proteins were determined by comparing mean values of
various dilutions of the mutant to the standard curve.

Biological activities and Immunizations.

Male C57BL/6 mice, 10 to 12-weeks old, were obtained from Harlan Sprague-Dawley, Inc. (Frederick Cancer Research and Development Center, Frederick, MD). The lethal effect of WT or mutant SEA was evaluated as described in Stiles et al. (1993) Infect. Immun. 61, 5333-5338. For immunizations, mice were given by interperitoneal (ip) injections either 2 or 10 μg of WT or mutant toxin in 100 μl of adjuvant (RIBI, Immunochem Research, Inc. Hamilton, MT or alum), or adjuvant only, and boosted (ip) at 2 and 4 weeks. Serum was collected from tail veins one week after the last immunization. Mice were challenged 2 25 weeks after the last injection with toxin and lipopolysaccharide (LPS, 150 μg) from E. coli 055:B5 serotype (Difco Laboratories, Detroit, MI). Challenge controls were adjuvant-immunized or non-immunized mice injected with both agents (100% lethality) or with either wild type toxin or LPS. No lethality was produced by these negative controls. Monkeys were immunized with the antigen in the right leg, caudal thigh muscles. Each received three intramuscular immunizations with a superantigen vaccine plus 35

adjuvant. Control monkeys received 0.5 ml total volume of adjuvant (Alhydrogel, Michigan Department of Public Health) and sterile PBS using the same techniques and equipment as the immunized monkeys. Immunizations were administered 28±2 days apart and consisted of 20 µg of the vaccine in adjuvant in a total volume of 0.5 ml. Immunizations were administered on day 0, 28±2, and 56±2 using a 23-27 ga 1/2-5/8" needle attached to a 1 ml tuberculin syringe into the caudal thigh.

Antibody assay.

Microtiter plates were coated with 1 µg/well of WT toxin in 100 µl of PBS (37°C, 2 h). After antigen 15 coating, the wells were blocked with 250 ul of casein 0.2% in PBS for 4 h at 37°C and then washed four times with PBS containing 0.2% Tween 20. Immune or nonimmune sera were diluted in PBS containing 0.02% casein and 100 µl of each dilution was added to 20 duplicate wells. After each well was washed four times, bound antibody was detected with horse radish peroxidase (Sigma Chemical Comp., St. Louis, MO) labelled goat anti-species specific IgG (37°C, 1 h), using O-phenylenediamine as the chromogen. Mean of duplicates OD (absorbance at 490 nm) of each treatment group was obtained and these data were compared on the basis of the inverse of the highest serum dilution that produced an OD reading four times above the negative control wells. For negative controls, 30 antigen or serum was omitted from the wells.

Superantigen binding and TCR subset analysis.

Cells from the mouse B-lymphoma line A20 (ATCC, Rockville, MD) $(2-4 \times 10^5 \text{ cells})$ were incubated (40 min

at 37°C) with WT or mutant toxin in Hanks balanced salt solution containing 0.5% bovine serum albumin (HBSS, USAMRIID). The cells were washed with HBSS and incubated with 5 µg of affinity-purified anti-toxin 5 antibody in HBSS (4°C, 45 min). Unbound antibody was removed and the bound antibody was detected with fluorescein isothiocyanate (FITC)-labelled, goat antirabbit IgG (Organon Teknika Corp., Durham, NC). Unbound antibody was removed and the cells were analyzed by with a FACSort flow cytometer (Becton Dikinson & Co. , Mountain View, CA).

For TCR subset analysis, splenic mononuclear cells were obtained from mice immunized with WT or mutant toxin. The mononuclear cells were incubated 15 (37°C) with WT toxin (100 ng/mL) for 5 days and then cultured in 85% RPMI-1640, 10% interleukin-2 supplement (Advanced Biotechnologies Inc., Columbia, MD) with 5% FBS for an additional 5 days. The T cells were washed twice and stained with anti-TCR 20 (Biosource, Camarillo, CA) or anti-Vß specific TCR (Biosource, Camarillo, CA) (45 min, 4°C). All cells analyzed were positive for T cell marker CD3+ and expressed the CD25 activation marker (data not shown). Controls were incubated with an isotype matched 25 antibody of irrelevant specificity. Unreacted antibody was removed, and the cells were incubated with an FITC-labelled, anti-mouse IgG (Organon Teknika Corp, Durham, NC) on ice for 30 min. The cells were washed and analyzed by flow cytometry (FACSort).

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LPS potentiation of SE toxicity in mice.

C57BL/6 or BALB/c mice weighing 18-20 g (Harlan Sprague Dawley, Inc., Frederick Cancer Research and Development Center, Frederick, MD) were each injected intraperitoneally (i.p.) with 200 µl of PBS containing

varying amounts of SEA, SEB, or SEC1, TSST-1, or SPEa followed 4 h later with 75 or 150 μg of LPS (200 $\mu l/i.p.$). Controls were each injected with either SE (30 mg) or LPS (150 mg). Animals were observed for 72 h after the LPS injection. Calculations of LD50 were done by Probit analysis using 95% fiducial limits (SAS Institute Inc., Cary, NC).

The biological effects of SEA and SEB were also tested in transgenic C57BL/6 mice (GenPharm

10 International, Mountain View, CA) deficient in MHC class I or II expression [Stiles et al. (1993) Infect. Immun. 61, 5333-5338], as described above, using a single dose of toxin (30 μg/mouse). Genetic homozygozity was confirmed by Southern analysis of parental tail DNA, using β2 microglobulin and MHC class II β DNA probes.

Detection of cytokines in serum.

Mice (n=18 per group) were injected with toxin

(10 μg), LPS (150 μg), or toxin plus LPS. Sera were
collected and pooled from three mice per group at each
time point (2, 4, 6, 8, 10, 22 h) after LPS injection.
Sera were collected at various time points following
toxin injection (-4 h, or 4h before LPS injection, for

data tabulation). Collection of LPS control sera
began at the time of injection (0 h).

Serum levels of TNF α and IL- α were detected by an enzyme linked immunosorbent assay (ELISA). TNF α was first captured by a monoclonal antibody against mouse TNF α (GIBCO-BRL, Grand Island, NY) and then incubated with rabbit anti-mouse TNF α antibody (Genzyme, Boston, MA). The ELISA plate was washed and peroxidase conjugate of anti-rabbit antibody (Boehringer Mannheim, Indianapolis, IN) added to the wells. After washing the plate and adding substrate (Kirkegaard and

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Perry, Gaithersburg, MD), TNFα concentrations were measured using the mean A450 reading of duplicate samples and a standard curve generated from redombinant mouse TNFa (GIBCO-BRL). Serum levels of 5 IL-1a were determined from the mean reading of duplicate samples with an ELISA kit that specifically detects murine IL-1a (Genzyme, Boston, MA). The standard error of the mean (SEM) for $TNF\alpha$ and $IL-l\alpha$ readings was +/- 5%.

Quantitation of IL-6 and IFNy were measured by bioassays [See et al.(1990) Infect. Immun. 58: 2392-2396]. An IL-6 dependent cell line, 7TD1 (kindly provided by T. Krakauer), was used in a proliferative assay with serial two-fold dilutions of serum samples assayed in triplicate. Proliferation of 7TD1 cells in a microtiter plate was measured by uptake of [3H]thymidine (1 µCi/well; Amersham, Arlington Heights, IL) and the activity of IL-6 from serum was compared to a recombinant mouse IL-6 standard (R and D Systems, Minneapolis, MN) as previously described [See et al. (1990) Infect. Immun. 58: 2392-2396]. The SEM of triplicate samples was +/- 10%.

IFNy was measured by the reduction of vesicular stomatitis virus (New Jersey strain) cytopathic effects on L929 cells, as previously described [Torre et al. (1993) J. Infect. Dis. 167, 762-765]. Briefly, serial two-fold dilutions of serum were made in duplicate and added to microtiter wells containing L929 cells (5 x 104/well). After incubating 24 h, 30 virus (5 x 10 PFU/well) was added and the cytopathic effects measured at 48 h by absorbance readings (570 nm) of reduced 3-[4, 5-dimethylthiazol-2-yl]-2,5 diphenyl tetrazolium bromide (Sigma). The activity of each serum sample was determined using recombinant

mouse IFNy as a standard (Biosource, Camarillo, CA).

The SEM of duplicate samples was +/- 5%.

EXAMPLE 1

5 <u>Molecular modelling and structural studies of</u> <u>staphylococcal and streptococcal superantigens:</u> <u>bacterial superantigens share common 3-dimensional</u> <u>structure.</u>

Comparison of amino acid sequences (Fig. 1) suggested that bacterial superantigens fall into 10 groups consisting of (1) SEA, SED and SEE, (2) SEB, staphylococcal enterotoxins C1-C3 (SEC1-3), the streptococcal pyrogenic exotoxins A (SPE-A) and C (SPE-C), (3) TSST-1 and (4) the exfoliative toxins 15 (ETA, ETB) and streptococcal pyrogenic exotoxin B (SPE-B), which are the most distant from the others in sequence. Although not available to the inventor when the inventions were first conceived and proof of principle was obtained, the x-ray crystallographic 20 structures of several bacterial superantigens are now known. Diverse superantigens, such as SEB and TSST-1, appear to have little sequence in common, yet they exhibit homologous protein folds composed largely of β strands [Prasad, G.S. et al. (1993) Biochemistry 32, 13761-13766; Acharya, R.K. et al. (1994) Nature 367, 25 94-97; Swaminathan, S. et al. (1992) Nature 359, 801-806] within two distinct domains. Differences between the proteins are located primarily in highly variable regions comprised of several surface loops, such as the disulfide-bonded loop which is absent from TSST-1 30 and at the amino terminus.

The X-ray crystal structures of SEB and TSST-1 complexed with HLA DR1 are known [Kim, J. et al. (1994) Science 266, 1870-1874; Jardetzky, T.S. et al.

(1994) Nature 368, 711-718] and this data was useful to fully explain our results concerning attenuation of the superantigens by site-specific mutagenesis. region of HLA DR1 that contacts SEB consists exclusively of $\boldsymbol{\alpha}$ subunit surfaces. The main regions of SEB involved are two conserved sites: a polar pocket derived from three β strands of the β barrel domain and a highly solvent-exposed hydrophobic reverse turn. The polar binding pocket of SEB 10 contains a glutamate and two tyrosines that accommodate Lys39 of the α subunit of HLA DR1, while the hydrophobic region consists of a leucine and flanking residues that make several contacts with the HLA DRa chain. The HLA DR1 binding sites of both 15 TSST-1 and SEB overlap significantly. The hydrophobic binding contacts of other SAg with the HLA $DR\alpha$ chain have been proposed [Ulrich et al. (1995) Nature, Struct. Biol. 2, 554-560] to be similar to those found in SEB and TSST-1. A motif consisting of a leucine in a reverse turn [Ulrich et al. (1995), supra] is conserved among bacterial superantigens and may provide the key determinant (hydrophobic or otherwise) for binding HLA-DR. However, TSST-1 does not have a highly charged residue in the polar pocket that interacts with Lys39 of the HLA $DR\alpha$ chain and uses an alternative conformational binding mode that allows TSST-1 to interact with HLA DR1 β -chain residues and the carboxy-terminal region of the antigenic peptide. Both SEA and SEE bind to the β subunit of DR by

Both SEA and SEE bind to the β subunit of DR by

30 means of a single zinc atom [Fraser, J.D. et al.
(1992) Proc. Natl. Acad. Sci. USA 89, 5507-5511]. The
amino-terminal domain of SEA interfaces with the HLA
DRα chain [Ulrich et al. (1995), supra], while SEA Cterminal domain residues His187, His225 and Asp227

35 form a zinc-coordination complex, likely with His-81

from the β chain of an adjoining HLA DR molecule. However, our results have shown that binding of superantigen to the HLA DRβ subunit does not directly stimulate T cells [Ulrich et al. (1995), supra] but increases the potential of the bound SEA to interact with the α chain of another HLA DR, thus increasing the biological potency.

EXAMPLE 2

Molecular modelling and structural studies of staphylococcal and streptococcal superantigens: A detailed protein structure analysis of SEB and SEA suggested that all bacterial superantigens have a common mechanism for binding MHC class II receptors.

15 A least-squares superimposition of the unbound molecules of modeled SEA and the crystal structure of SEB, aligned according to their structurally conserved α-helical and β-strand regions, exhibited a global folding pattern which is very similar. Differences between the two structures are calculated to be 20 located primarily in loops of low sequence homologies, with the largest positional deviations occurring between structurally conserved regions of residues 18-20, 30-32, 173-181, 191-194, and the cysteine-loop region (90-111). Only one of these regions in SEB 25 makes significant contact (residue Y94 in particular) with the HLA-DR1 molecule [Jardetzky, T.S. et al. (1994) Nature 368, 711-718].

The binding interface between SEB and HLA-DR1 consists principally of two structurally conserved surfaces located in the N-terminal domain: a polar binding pocket derived from three ß-strand elements of the ß-barrel domain and a hydrophobic reverse turn. The binding pocket of SEB contains residues E67, Y89

and Y115, and binds K39 of the DRa subunit. For SEA, the binding interface with the DR molecule is modeled to contain a similar binding pocket consisting of residues D70, Y92 and Y108. Mutation of residue Y89 5 in SEB or Y92 in SEA to alanine (Fig. 2) resulted in 100-fold reduction in DR1 binding. The substitution of alanine for Y89 in SEB and Y92 in SEA eliminates the hydrogen bond with K39 and disrupts packing interactions with adjacent protein residues. Modeling of the SEA mutant Y92A predicts an increase in 10 solvent-accessible surface area for Y108 by a factor of two greater than the wild-type structure, allowing the formation of a hydrogen bond to the carboxylate group of D70 and thus disrupting key anchoring and 15 recognition points for HLA-DR1. This effect is expected to be somewhat less in SEB due to the longer side chain at E67. Substitution of SEB Y115 with alanine also resulted in 100-fold reduction of binding. In contrast, the same replacement of Y108 in SEA yielded little to no change in DR1 binding (Fig. 20 2a), suggesting the primary importance of SEA residues Y92 and D70 for stabilizing interactions with K39. The K39 side chain of DR α forms a strong ion-pair interaction with the SEB E67 carboxylate group and 25 hydrogen bonds with the hydroxyl groups of Y89 and Y115. Substitution of SEB E67 by glutamine reduced binding affinity by 100-fold (Fig. 2), reflecting the replacement of the strong ionic bond with a weaker hydrogen bond. To optimize ion-pair interactions of the analogous SEA site, the shorter carboxylate side chain of D70 is predicted to shift K39 of $DR\alpha$, weakening interactions with SEA Y108. substitution of alanine for SEA Y108 is thus more easily accommodated than the homologous substitution of SEB Y115, without loss in DR1 binding.

Comparisons of the polar pocket with other bacterial superantigens were then made. SEC1-3 and SPE-A have conserved the critical DR1 bindinginterface residues (Fig. 1), and share with SEB and 5 SEA secondary structural elements of the DR1-binding surfaces. Asparagine in SED (N70) replaces the acidic side chain present in SEA, SEB, SPE-A and SEC1-3. Accordingly, for SED the salt bridge of the polar pocket is likely to be replaced by a hydrogen bond. 10 Overall DR1 affinities for SED and SEA appeared to be equivalent (Fig. 2b), indicating that other interactions may compensate for the absence in SED of the ion-pair found in the other superantigens. For the case of TSST-1, mutating DRa residues K39 to serine or M36 to isoleucine has been shown to greatly 15 reduce binding [Panina-Bordignon et al. (1992) J. Exp. Med. 176: 1779-1784]. Although primarily hydrophobic, the critical TSST-1 structural elements are conserved with the SEA and SEB polar binding 20 pocket. SEB residues Y89 and Y115 are homologous to T69 and I85 in TSST-1, respectively, and SEB E67 is replaced by I46. These TSST-1 residues are positioned in a conserved &-barrel domain found in both SEB and SEA. However, the TSST-1 site lacks polarity equivalent to SEB/SEA, and hydrogen bonding with the 25 hydroxyl of TSST-1 residue T69 would require that $DR\alpha$ K39 extend 5 Å into the pocket. TSST-1 binding utilizes an alternative strategy [Kim et al. (1994) Science 266: 1870-1874] consisting of hydrophobic 30 contacts centered around residue I46, and potential ionic or hydrogen bonds bridging DRa residues E71 and K67 to R34 and D27, respectively, of TSST-1.

The hydrophobic region of the binding interface between SEB and the HLA-DR1 molecule consists of SEB residues 44-47, located in a large reverse turn

connecting ß-strands 1 and 2 of SEB. These residues appear to make strong electrostatic interactions with DRa through their backbone atoms. The mutation of L45 to an arginine reduced overall HLA-DR1 binding greater 5 than 100-fold (Fig. 2b), attributable to the less energetically favorable insertion of a highly charged residue into a hydrophobic depression on the DR1 molecule. The modeled DR1-SEA complex presents similar interactions with the SEA backbone atoms, with 10 the exception of a glutamine (Q49) replacing SEB Y46. Mutation of L48 to glycine in SEA (homologous to L45 of SEB) has been reported to decrease T-cell responses. SEB L45 and the comparable L30 of TSST-1 are the most extensively buried residues in the DR1 15 interface. The leucine is conserved among the bacterial superantigens (Fig. 3) and may provide the necessary hydrophobic structural element for surface complementarity with DR1, consistent with the mutagenesis data for SEB and SEA.

The inventor has performed similar structure and function studies with TSST-1, SEC1 and SPE-A.

EXAMPLE 3

Molecular modelling and structural studies of

staphylococcal and streptococcal superantigens: Some
interactions of bacterial superantigens with MHC class
II receptors are not conserved but are less important
than the hydrophobic loop and polar pocket binding
sites.

In determining the overall affinity of the superantigen for DR1, a contributory role is played by structural variations around the common binding motifs. A short, variable structured, disulfide-bonded loop is found in SEA and a homologous longer

loop in SEB. The SEB residue Y94, contained within this loop, forms hydrophobic interactions with L60 and A61 of the DRa subunit. Replacement of Y94 with alanine partially inhibits DR1 binding (Fig. 2a,b). An alanine is found in SEA (A97) and SEE at the position equivalent to SEB Y94, and mutating this residue in SEA to tyrosine results in disrupted instead of stabilized interactions with DR1 (Fig. 2a). Although the disulfide loops differ in structure between SEA and SEB, A97 apparently contributes to the 10 DRa binding interface in a manner similar to Y94 of SEB. Because TSST-1 lacks a disulfide loop, similar contacts with DRa are replaced by interactions with β strands of TSST-1. In a like manner, the absence of a salt bridge between the residues K39 of DR α and E67 of SED is apparently compensated for by stabilizing interactions occurring outside of the otherwise conserved dominant binding surfaces (Fig. 2a).

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EXAMPLE 4

Molecular modelling and structural studies of staphylococcal and streptococcal superantigens:
Superantigen interactions with T-cell antigen receptors.

25 The amino acid residues in contact with TCR are located in regions of high sequence variability, presenting a unique surface for interaction with the TCR. Residues implicated in TCR interactions by mutagenesis of SEA and SEB reside in variable loop regions, while TSST-1 mutants that affect TCR binding are mainly located in an α helix [Acharya, R.K. et al. (1994) Nature 367, 94-97; Kim, J. et al. (1994) Science 266, 1870-1874]. Specifically, mutations that diminish T-cell receptor recognition of SEB include

residues N23, Y61, and the homologous SEA N25 or Y64 (Fig. 2c). SEA residues S206 and N207 also control T-cell responses [Hudson, et al. (1992) J. Exp. Med. 177: 175-184]. Mutants of the polar binding pocket, 5 SEA Y92A and SEB Y89A, equivalently reduced T-cell responses (Fig. 2c), reflecting the observed decreases in DR1-binding (Fig. 2a, b). While supporting reduced T-cell responses, mutants SEA Y64A and SEB Y61A retained normal affinities for DR1 (Fig. 2a-c).

EXAMPLE 5

Animal models for determining biological activity of bacterial superantigens: Mouse.

When compared to primates, mice are not very 15 susceptible to the toxic effects of SE, and we therefore sought to increase sensitivity with a potentiating dose of lipopolysaccharide (LPS) from Gram-negative bacteria [Stiles et al. (1993) Infect. Immun. 61, 5333-5338]. There was no apparent effect 20 in control animals injected with any of the SE (up to 30 μ g/mouse) or LPS (150 μ g/mouse) alone (Table 1). Incremental injections of LPS were also not lethal, when given in amounts up to 250 μg/mouse (data not shown). However, mice died between 24-48 h after SE 25 and LPS were given to the same animal (Table 1). SEA was much more toxic than either SEB or SEC1 and the calculated LD50 (µg toxin/kg) of SEA, SEB, and SEC1 with 95% fiducial limits was 18.5 (6.5, 38.5), 789.0 (582.5, 1044.5), and 369.0 (197.5, 676.0), 30 respectively.

TABLE 1. Titration of SEA.SEB, and SEC, in the C57BL/6 mouse lethality assay

5 % Lethality (no. of mice tested) with Stimulus the following dose of SE, in micrograms/mouse^b:

10		30	10	1	0.1
10	SEA + LPS	93 (15)b	85 (20)	80 (15)	20(10)
	SEB + LPS	80(15)	27 (15)	0(15)	0(15)
	SEC ₁ + LPS	80(10)	60(10)	10(10)	0(10)

15 *LPS was injected into each mouse (150ug) 4 h after the SE injection. Control mice injected with 150 ug of LPS (n=20) or 30 ug of SEA, SEB, or SEC1 (n=10) survived.

 $^{\mathtt{b}} Results$ are from a combination of separate experiments with five mice per experiment.

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The role of MHC class I and class II molecules in SE toxicity, potentiated by LPS, was addressed by using transgenic, MHC-deficient mice (Table 2). Class II-deficient animals were unaffected by a dose of SE (30 μ g) plus LPS (150 μ g) that was lethal for 93% of wild-type and 30% of class I-deficient mice. Mononuclear cells from class II-deficient animals were not able to present SEA, as measured by proliferative responses. MHC class I-deficient cells were functional in supporting T-cell proliferation, but at levels <30% of the proliferative response supported by MHC-wild-type presenting cells (Table 3). Cell surface expression levels were normal, when compared to nontransgenic C57BL/6, for Ab in class I-deficient mice, and K^b/D^b in class II-deficient mice. The Tcell responses of MHC class I- or class II-deficient

mice were essentially equivalent to wild-type when SEA

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was presented by mononuclear cells expressing both class I and II molecules (Table 3).

TABLE 2. Lethality of SEA and SEB in C57BL/6 mice

5 lacking MHC class I or class II

% Lethality (no. of mice tested) with
Stimulus* the following MHC class phenotype

		I.II.	1,11.	1,11,		
SI	A + LPS	30(10)	0(5)	93 (15)		
SE	EA + LPS	NDp	0(5)	80(15)		
SE	EA only	0(2)	0(2)	0(2)		
SE	B only	NDp	0(2)	0(2)		
LF	S only	0(5)	0(5)	0(5)		

^{*} Mice were injected with 30 ug of SEA or SEB and, 4h later, with 150 ug of LPS, as indicated. Control mice were injected with only SEA, SEB, or LPS.

b ND, not determined.

Table 3. Mouse T-cell responses to SEA are MHC class II-dependent

T-cell responses

	T-cell/APC source	0.1 µg/ml SEA	1 μg/ml SEA
30	Wild-type C57/BL6 mouse/autologous	430,000 cpm ²	700,000 cpm
35	MHC class I knock-out C57/BL6 mouse/autologo	117,000 cpm us	167,000 cpm
40	MHC class II knock-out C57/BL6 mouse/autologo	8,000 cpm	33,000 cpm
10	Wild-type C57/BL6 mouse/wild-type	305,000 cpm	307,000 cpm
45	MHC class I knock-out	420,000 cpm	445,000 cpm

C57/BL6 mouse/wild-type

MHC class II knock-out C57/BL6

310,000 cpm

322,000 cpm

5 mouse/wild-type

¹Cultures of mononuclear cells derived from mouse spleens, cultured for 3 d with the indicated amount of SEA.

10 ²Data represent the mean of triplicate determinations (<10 SEM) of [³H)thymidine incorporation.

³Antigen presenting cells (APC) were isolated from spleens of the indicated mouse strain and added to cultures.

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The serum levels of TNFα, IL-1α, IL-6, and IFNγ in mice injected with SEA, LPS, or SEA plus LPS were measured at various times following injection (Fig. 4). Compared to mice injected with either SEA or LPS alone, the serum levels of TNFα, IL-6, and IFNγ had increased 5-, 10-, and 15-fold, respectively, in animals given SEA plus LPS. SEA alone did not elicit any detectable increase of TNFα, IL-6, or IFNγ above background. In contrast to the other cytokines, IL-1α levels in mice injected with SEA plus LPS resulted in a simple additive effect.

Serum levels of TNFα, IL-6, and IFNγ were maximal 2-4 h after the LPS injection, but returned to normal by 10 h. The concentration of IL-1α in mice given SEA plus LPS had also peaked 2 h after the LPS injection, but stayed above background for the remaining determinations. Levels of IL-1α in mice given only LPS or SEA peaked at 4 and 6 h, respectively. Unlike profiles for other cytokines, the highest amount of IL-1α in mice injected with SEA and LPS corresponded to the peak stimulated by SEA, but not LPS.

This animal model was used in various stages of developing the inventions, as a means of assessing the physiological activity of mutated superantigens.

Control animals survived the maximum dose of either SE or LPS, while mice receiving both agents died. Wildtype SEA was 43-fold more potent than SEB and 20-fold more potent than SEC1. By using BALB/c mice the toxicity of SEB was 10-20 fold higher. These data confirmed that the toxicity of SE was mainly exerted through a mechanism dependent on expression of MHC class II molecules and was linked to stimulated cytokine release. Thus this was a relevant preclinical model that could be used to predict human responses.

EXAMPLE 6

Animal models for determining biological activity

of bacterial superantigens: Rhesus monkey

The physiological responses of the rhesus monkey to bacterial superantigens is probably identical to humans, with the exception of sensitivity [Bayari and Ulrich (1995) Clin. Immunol. Immunopath. 76:2481. 20 Generally SEB intoxicated monkeys developed gastrointestinal signs within 24 hours post-exposure. Clinical signs were mastication, anorexia, emesis and diarrhea. Following mild, brief, self-limiting gastrointestinal signs, monkeys had a variable period 25 of up to 40 hours of clinical improvement. At approximately 48 hours post-exposure, intoxicated monkeys generally had an abrupt onset of rapidly progressive lethargy, dyspnea, and facial pallor. given a lethal dose, death occurs within four hours of 30 onset of symptoms. Only SEB has been used in challenges of rhesus monkeys to determine physiological/pathological effects. Human responses to bacterial superantigens are characterized by a rapid drop in blood pressure, elevated temperature,

and multiple organ failure-the classical toxic shock syndrome (TSS). However, the respiratory route of exposure may involve some unique mechanisms. The profound hypotension characteristic of TSS is not observed, and respiratory involvement is rapid, unlike TSS. Fever, prominent after aerosol exposure, is generally not observed in cases of SEB ingestion.

EXAMPLE 7

10 <u>Targeting receptor interactions to develop</u> vaccines.

The SEA mutants Y92A, with reduced DR1 binding, and Y64A, with reduced TCR interactions, and K14E with wild-type (control) activity were used to determine 15 the correct receptor to target for vaccine development. The binding of WT or mutant SEA was evaluated with the MHC class II expressing murine Bcell lymphoma cell line A20 (Table 4). The binding affinity of WT SEA to mouse MHC class II (H-2d) 20 molecules was lower than that observed with human MHC class II expressing cells , reflecting the reduced toxicity that bacterial SAgs exert in mice. WT SEA, Y64A and K14E all had the same relative affinity to mouse MHC class II molecules. Similar to the results 25 obtained with human MHC class II molecules, the Y92A mutant exhibited substantially reduced binding to A20 cells (Table 4).

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Table 4. Biological activity of superantigen vaccines .

5	toxin T-cel	ll anergy¹	MHC classII binding ²	T-cell response
	SEA wild type	++++	+++	+++
10	TCR attenuated Y64A	+	+++	+/-
	MHC attenuated Y92A	-	+/-	. +/-
15	Control K14E	++++	+++	+++

¹Based on attenuation of T-cell response to wild-type SEA in mice immunized with the mutant or wild-type SEA.

25 The effect of WT SEA or site-specific SEA mutants on splenic mononuclear cells obtained from nonimmunized C57BL/6 (H-2b) mice is summarized in Table 4. Both WT SEA and the control mutant K14E were potent T cell activators, effective at minimal 30 concentrations of 10 to 100 pg/mL. However, T-cell responses to Y92A were reduced at least 100-fold, compared to SEA wild type, while Y64A-stimulated responses were slightly higher than Y92A. These results confirmed that attenuation of superantigen binding to either MHC class II or TCR molecules 35 resulted in dramatically reduced mouse T-cell proliferation. These results may indicate that the altered toxin may compete with wild type toxin for TCR binding.

40 SEA WT (10 LD50), site-specific SEA mutants (10 μg/mouse each) or LPS (150 μg/mice) injected alone were nonlethal to mice (Table 5). However, combining

 $^{^{2}\}mbox{Binding}$ to the mouse MHC class II+ A20 cells, measured by flow cytometry

LPS with either WT SEA or mutant K14E resulted in 100% lethality. For those mice receiving both LPS and WT or K14E SEA, 80% were dead by 24 h and 100% by 48 h. In contrast, 100% of Y92A and 80% of Y64A injected

5 mice (coadministered with LPS) survived. The average time to death for the 20% of mice that did not survive Y64A injection occurred at 48 to 72 h. These in vivo data correlated well with the results obtained with the lymphocyte cultures. It was concluded that the observed attenuation of toxicity in mice was a direct result of the reduced T-cell proliferation.

Table 5. Biologic effect of wild type (WT) staphylococcal enterotoxin A (SEA) and SEA mutants.

Protein	No. live/total
WT	0/10
K14E	0/10
Y64A	8/10
Y92A	10/10

NOTE. Mice were given 10 LD_{50} (10ug) of WT or mutant SEA. Lipopolysaccharide (150 ug/mouse) was injected 3 h later.

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Having established that attenuation of receptor binding resulted in reduced toxicity, we next examined the immunogenicity of the SEA mutants. Mice were immunized with WT or mutant SEA. Control mice received adjuvant only or were left untreated. One week before challenge with WT SEA, mice were bled and serum antibody titers were determined for each group (Table 6). Mice immunized with the 2 µg of Y64A or Y92A had serum antibody titers of 1:5000 and 1:1000, respectively. Immunization with 2 µg of WT SEA or

control mutant resulted in titers of 1:5,000 and 1:10,000, respectively. The highest immunizing dose (10 μ g/mouse) was most effective for all animals, resulting in antibody titers which were greater than 1:10,000. All mice were challenged with 10 LD50 of WT SEA (potentiated with LPS). The survival data correlated well with the levels of serum antibodies in immunized mice. All mice that were vaccinated with 10 μg of Y64A or Y92A, survived the lethal challenge dose of WT SEA. Slightly less protection was afforded by 10 the lower vaccination dose of mutant Y64A or Y92A. All mice immunized with both doses of WT SEA survived the lethal challenge with WT potentiated with LPS. Mice immunized with mutant K14E exhibited survivals of 100% and 80% for high and low vaccination doses, respectively. All nonimmunized or control mice that were vaccinated with adjuvant alone died when challenged with WT SEA and a potentiating dose of LPS.

20 Table 6. Mice immunized with attenuated forms of staphylococcal enterotoxin A (SEA) produce high titers of neutralizing antibody.

Immunizing agent	Dose (ug/mouse)	Anti-SEA antibody titer*	No. live/tota
WT	2	10,000-50,000	10/10
	10	10,000-50,000	10/10
K14E	2	5,000-10,000	8/10
	10	10,000-50,000	10/10
Y64A	2	5,000-10,000	6/10
	10	10,000-50,000	10/10
Y92A	2	1,000-5,000	2/10
	10	10,000-50,000	10/10
Adjuvant		50-100	0/10

NOTE. Mice were given 10 LD_{50} of wild type (WT) SEA challenge followed by potentiating dose of lipopolysaccharide (150 ug/mouse) 3 h later.

*Reciprocal of serum dilution resulting in optical density reading four times above negative controls (wells containing either no SEA or no primary antibody).

EXAMPLE 8

Immune recognition of SAg mutants.

10 Bacterial SAgs induce clonal anergy of specific subsets of T cells in mice. It was possible that the loss of sensitivity to WT SEA among the mice vaccinated with the attenuated mutant forms represented a state of specific non-responsiveness 15 instead of specific immunity. To address this issue, lymphocyte responses to SEA WT were measured with splenic mononuclear cells collected 2 weeks after the third immunization. As expected, lymphocytes from mice that were immunized with WT SEA or control SEA mutant showed little to no proliferation when 20 incubated with the WT SAg. In contrast, lymphocytes obtained from control mice or those immunized with either Y64A or Y92A all responded vigorously to the WT SEA (Fig. 5). The TCRs used by T cells from the SEAvaccinated mice were then characterized by flow cytometry. T cells from immunized or control mice were incubated with WT SEA in culture for 7 days, followed by a 5 day expansion in IL-2 containing medium. Distinct populations of activated TCR VB11 positive cells were observed with T cells from mice immunized with Y92A and Y64A, representing 48% and 40% of T cells, respectively. However, VB11 expressing cells obtained from SEA WT or K14E immunized mice were about 1% and 6% of the total T-cell population, 35 respectively, suggesting that this subset was

nonresponsive to restimulation with the WT SAg. T cells bearing Vß 17a, 3, 7, and 10b were unchanged for all mice. It was apparent that T-cell responses to both the TCR and MHC class II binding-attenuated SEA mutants were similar to each other, but differed from responses to control or WT molecules. These results suggested that an alternative, perhaps conventional antigen processing mechanism was functioning in presentation of the SAg mutants Y64A and Y92A.

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EXAMPLE 9

Rhesus monkey immunizations with monovalent vaccines.

The SEA vaccine L48R, Y89A, D70R (A489270) and 15 SEB vaccine Y89A, Y94A, L45R (B899445) were used to immunize rhesus monkeys. The animals received a total of three i.m. injections (10-20 μ g/animal), given at monthly intervals. Rhesus monkeys that were injected with these vaccines had no detectable increase of 20 serum cytokines and no apparent toxicity. The serological response of animals vaccinated with three doses of formalin-treated SEB toxoid (100 μg/injection) gave results comparable to one or two injections with B899445 (Table 7), suggesting that the recombinant vaccines were very immunogenic. Immunized 25 rhesus monkeys survived a lethal challenge with >10 LD50 of wild-type SEB (Table 7, 8). Collectively, these results suggest that the engineered SEB vaccine is safe, highly antigenic and effective at protecting the immunized individual from lethal aerosol exposure 30 to SEB.

Table 7. Rhesus monkey antibody responses to vaccine B899445; One injection of B899445 outperforms three injections of SEB toxoid

5	Vaccine ¹ /animal #	Antibody response ²	%Inhibition of T-cell response ³	Survival SEB >20 x LD50 challenge
10	pre'immune sera /pooled	0.161	5	dead
	toxoid/1	0.839	0	dead
15	toxoid/2	0.893	34	live
13	toxoid/3	1.308	57	live
	toxoid/4	1.447	55	live
20	B899445/1	1.788	69	live
	B899445/2	0.78	49	live

 1Rhesus monkeys were immunized with one dose (20 μg injection) of B899445 vaccine or three doses of formalin-treated SEB toxoid (100 $\mu g/injection)$ one month apart; both used Alum adjuvants.

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²Sera were collected one month after the final injection. Antibody responses were determined by ELISA and the results are shown as mean optical densities of triplicate wells (± SEM).

35 Rhesus monkey T cells, obtained from an untreated animal, were preincubated with diluted (1:70) serum from immunized monkeys and then cultured with wild type SEB. Data are shown as % of T cell responses, where serum of rhesus monkey injected with adjuvant only represented the 100% of response to wild type SEB.

 $\ensuremath{^4\mathrm{Rhesus}}$ monkeys were challenged by aerosol exposure and monitered for four days.

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Table 8. Engineered staphylococcal enterotoxin B vaccine efficacy in rhesus monkeys

5	Treatment'	Antibody titer	Immune protection'	
	Vaccine with adjuvant	>10,000	100%	_
10	Adjuvant only	<50	0%	

¹Rhesus monkeys (n=10) were injected i.m. with 10 μ g of SEB vaccine with Alhydrogel adjuvant. A total of 3 immunizations, 1 month apart were given. Controls (n=2) received only Alhydrogel.

²Serum dilution resulting in optical density readings of four times above the negative control, consisting of no SEB or serum added to the wells.

³Immunized and control rhesus monkeys were challenged with >10 LD50 of wild-type staphylococcal enterotoxin B as an aerosol.

25 Serum from monkeys that were immunized with the genetically attenuated vaccine inhibited T-lymphocyte responses to wild type SEB (Table 7) similarly or better than monkeys that received the SEB toxoid. Collectively, these results suggest that the recombinant SAg vaccines are safe, highly antigenic, and induce protective immunity.

Serum from B899445 immunized rhesus monkeys blocked human lymphocyte responses to wild-type superantigen when tested in ex vivo cultures (Table 7). These data again showed that the second and third injections of vaccine were approximately equivalent in stimulating neutralizing antibody responses. Normal T-cell responses to several superantigens, including the wild-type protein, were observed in immunized animals, indicating that no specific or generalized anergy occurred (Fig. 6).

EXAMPLE 10

A. <u>Multivalent superantigen vaccines: Rhesus</u> monkey immunizations.

Rhesus monkeys were immunized with a combined vaccine consisting of B899445 and A489270. Following the third injection, antibody recognition of wild-type bacterial superantigens was examined (Fig.7). High titers of anti-SEB, SEC1 and SEA antibodies were evident.

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B.Mouse immunizations.

Mice (BALB/c) were immunized with a combined vaccine consisting of SEA, SEB, SEC1 and TSST-1 (all wild-type). The antibody responses against each 15 individual superantigen were assessed (Table 9). Antibodies were induced against each of the component antigens, providing sufficient levels to protect the mice from a lethal challenge of superantigen, potentiated with LPS. Although not shown in the 20 Table, antibody responses against SPE-A were also observed. Mice were also immunized with individual superantigens and antibody responses against other superantigens were measured (Table 10). Each individual immunogen induced partial or complete 25 protective antibody responses against all other superantigens tested.

TABLE 9. Superantigen cross-reactivity of antibodies from mice immunized with individual bacterial superantigens

Immunizing ¹ Toxin	Challenging ² Toxin	ELISA ³ Titer	Neutralizing ⁴ Antibody
SEA	SEA	>1/25,000	100%
SEA	SEB	>1/25,000	100%
SEA	SEC1	>1/25,000	100%
SEA	TSST1	>1/10,000	100%
SEB	SEB	>1/25,000	100%
SEB	SEA	>1/10,000	100%
SEB	SEC1	>1/2,500	100%
SEB	TSST1	>1/10,000	100%
SEC1	SEC1	>1/10,000	100%
SEC1	SEA	>1/10,000	100%
SEC1	SEB	>1/25,000	100%
SEC1	TSST1	>1/10,000	100%
TSST1	TSST1	<1/10,000	100%
TSST1	SEA	<1/1,000	50%
rsst1	SEB	<1/1,000	40%
rsst1	SEC1	<1/1,000	40%

²⁵ Three injections with 20 ug of antigen (BALB/c mice).

 $^{^2} LPS\mbox{-potentiated}$ challenge with 10 $LD_{50}s$ of superantigen.

³ELISA antibody response against an individual superantigen.

 $[\]ensuremath{^4\mathrm{Percent}}$ mice surviving an LPS-potentiated challenge 30 $\ensuremath{\text{(n=10)}}$.

Table 10. Multivalent superantigen vaccine. Mouse immune responses.

5	Immu to 	nizing n'	Ch	allenging toxin'	Antibody Titer'	% survival
	SE-A,	B, C1,	TSST-1	all	N/A	100%
10	11	•		SEA	>25,000	100%
		n		SEB	>25,000	100%
15	н	n		SEC1	>25,000	100%
.5	**	n		TSST-1	>6,400	100%

^{20 &}lt;sup>1</sup>Total of three injections, two weeks apart, in RIBI adjuvant.

'Measured by ELISA.

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EXAMPLE 11

Design of altered TSST-1 toxin vaccine, TST30.

A comprehensive study of the relationships of TSST-1 protein structure to receptor binding were undertaken to provide insight into the design of the 30 vaccine TST30. We have discovered that TSST-1 interactions with the human MHC class II receptor, HLA-DR, are relatively weak and can be disrupted by altering only a single critical amino acid residue of 35 the toxin. Site-directed mutagenesis of a gene encoding the toxin and expression of the new protein product in E. coli were then used to test the design of the vaccine. The TSST-1 gene used was contained within a fragment of DNA isolated by BglI restriction enzyme digestion of the gene isolated from a toxigenic 40 strain of Staphylococcus aureus (AB259; Kreiswirth and Novick (1987) Mol. Gen. Genet. 208, 84-87). sequence of this gene is identical to all currently

²>10 X LD50, potentiated with E. coli lipopolysaccharide.

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known TSST-1 isolates of human origin. The wild-type TSST-1 gene can be readily cloned from a number of clinical S. aureus isolates. The DNA fragment containing the TSST-1 gene was isolated by agarose gel electrophoresis and ligated inot the prokaryotic expression vector pSE380 (Invitrogen Corp.). The DNA clone consisted of sequences encoding the leader peptide and the full length of the mature TSST-1 protein. This engineered vaccine is currently being 10 evaluated to determine mouse and human T-cell reactivities in vitro, and lethality in mice. The TST30 vaccine consists of the following mutation introduced into the toxin molecule: leucine at amino acid residue 30 changed to arginine. Two other mutations, namely Asp27 to Ala and Ile46 to Ala have 15 also been designed. The final vaccine may incorporate one or both of these additional mutations.

The binding interface between TSST-1 and HLA-DR consists of a large relatively flat surface located in the N-terminal domain. Leucine 30 protrudes from a reverse turn on the surface of TSST-1 and forms the major hydrophobic contact with the HLA-DR receptor molecule. Mutation of the single residue leucine 30 in TSST-1 to the charged amino acid side chain of arginine is predicted to disrupt this major contact with the receptor molecule, resulting in a significant reduction in DR1 binding. This mutant molecule should therefore have lost the toxin attributes of the wild-type molecule.

TST30 was expressed as a recombinant protein in E.coli, as either a periplasmically secreted protein or as a cytoplasmic product. Purification was achieved by immunoaffinity chromatography or preparative isoelectric focusing after an initial ion-exchange CM-Sepharose enrichment step. The method of

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purification was not critical to the performance of the vaccine. Lipopolysaccharide contaminants, resulting from expression in a Gram-negative bacterium, were readily removed (as determined by limulus assay) using a variety of standard methods. The final purified vaccine is not toxic to mice at levels equivalent to 10 LD₅₀ of the native TSST-1. No indicators of toxicity were found in surrogate assays of human T-cell stimulation.

Conclusive vaccine studies demonstrating that TST30 is highly antigenic and induces protective immunity are in progress in a mouse animal model. Mouse lethality is achieved at less than 1 ug/animal when a potentiating signal like lipopolysaccharide from Gram-negative bacteria (LPS) is provided. When coadministered with LPS, wild-type TSST-1 is 100% lethal to mice (10 LD₅₀). Mice receive three injections (two weeks between injections) of 20 ug/mouse in alhydrogel and protection against the lethal effects of 10 LD₅₀ of TSST-1 are assessed.

EXAMPLE 12

Design of altered SPEA toxin vaccine, SPEa42

The SPEa interactions with human MHC class II receptor, HLA-DR, are relatively weak and can be disrupted by altering only a single critical amino acid residue of the toxin. Site-directed mutagenesis of a gene encoding the toxin and expression of the new protein product in E.coli were then used to test the design of the vaccine. The SPEa gene used was clone from a SPEa-toxigenic strain of Streptococcus by using specific DNA oligonucleotide primers and the polymerase chain reaction method. The sequence of this gene is identicla to SPEa isolates of human origin known within the public domain. The DNA

fragment containing the SPEa gene was isolated by agarose gel electrophoresis and ligated into a prokaryotic expression vector (pETx or pSE380). The DNA clone consisted of sequences encoding the leader peptide and the full length of the mature SPEa protein or SPEa42 without a leader sequence. We recognize that there are additional ways to express or produce the mature SPEa vaccine. The SPEa vaccine consists of the following mutation introduced into the toxin molecule: leucine at amino acid residue 42 changed to arginine.

The binding interface between SPEa and HLA-DR is predicted to consist of contacts located in the N-terminal domain that are conserved with other

15 bacterial superantigens. Leucine 42 of SPEa is predicted to protrude from a reverse turn on the surface of SPEa and form a major hydrophobic contact with the HLA-DR receptor molecule. Mutation of the single residue leucine 42 in SPEa to the charged amino acid side chain of arginine is predicted to disrupt this major contact with the receptor molecule, resulting in a significant reduction in DR1 binding. This mutant molecule should therefore have lost the toxin attributes of the wild-type molecule.

SPEa42 was expressed as a recombinant protein in E.coli, as either a periplasmically secreted protein or as a cytoplasmic product. Purification was achieved by immunoaffinity chromatography or preparative isoelectric focusing after an initial ion-exchange CM-Sepharose enrichment step. The method of purification was not critical to the performance of the vaccine. Lipopolysaccharide contaminants, resulting from expression in a Gram-negative bacterium, were readily removed (as determined by limulus assay) using a variety of standard methods.

The final purified vaccine is not toxic to mice at levels equivalent to 10 LD_{50} of the native TSST-1. No indicators of toxicity were found in surrogate assays of human T-cell stimulation.

Conclusive vaccine studies demonstrating that SPEa42 is highly antigenic and induces protective immunity are in progress in a mouse animal model. Mouse lethality is achieved at less than 1 ug/animal when a potentiating signal like lipopolysaccharide from Gram-negative bacteria (LPS) is provided. When coadministered with LPS, wild-type SPEa is 100% lethal to mice (10 LD₅₀). Mice receive three injections (two weeks between injections) of 20 ug/mouse in alhydrogel and protection against the lethal effects of 10 LD₅₀ of SPEa are assessed

EXAMPLE 13

Design of altered superantigen toxin vaccine. SEC45

20 For Staphylococcal enterotoxin C1 (SEC1), the leucine at position 45 was changed to lysine (SEC45). This mutation is anticipated to prevent SEC1 from interacting with the MHC class II receptor by sterically blocking the hydrophobic loop (centered around leucine 45) from binding to the alpha chain of 25 the receptor. SEC1 is more closely homologous to SEB than SEA or the other superantigen toxins. The presence of zinc in SEC1 may impart additional binding characteristics that allow, in some cases, this superantigen toxin to bind to T-cell antigen receptors without the required MHC class II molecule interactions. To circumvent the binding to T-cell antigen receptors, mutations of SEC1 residues N23

(changed to alanine), V91 (changed to lysine) are being performed.

SEQUENCE LISTING

5	(1) GENERAL INFORMATION: (i) APPLICANT: Robert G. Ulrich, Mark A. Olson Sina Bavari
10	(ii) TITLE OF INVENTION: Bacterial Superantigen Vaccines
	(iii) NUMBER OF SEQUENCES:16
15	(iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: John Moran (B) STREET: US Army MRMC -504 Scott Street MCMR-JA (John Moran-Patent Atty)
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25	 (v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: Apple Macintosh (C) OPERATING SYSTEM: Macintosh 7.5 (D) SOFTWARE: Microsoft Word 6.0
30	 (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: 08/882,431 (B) FILING DATE: June 25, 1997 (C) CLASSIFICATION:
35	(Vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE:
40	<pre>(viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Moran, John (B) REGISTRATION NUMBER: 26,313 (C) REFERENCE/DOCKET NUMBER:</pre>
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	(2) INFORMATION FOR SEQUENCE ID NO:1: (i) SEQUENCE CHARACTERISTICS:
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	(C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: DNA	
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		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
10		Met Tyr Lys Arg Leu Phe Ile Ser His Val 5 10
15		Ile Leu Ile Phe Ala Leu Ile Leu Val Ile 15 20
13		Ser Thr Pro Asn Val Leu Ala Glu Ser Gln 25 30
20		Pro Asp Pro Lys Pro Asp Glu Leu His Lys 35 40
		Ser Ser Lys Phe Thr Gly Leu Met Glu Asp 45 50
25		Met Lys Val Leu Tyr Asp Asp Asn His Val 55 60
30		Ser Ala Ile Asn Val Lys Ser Ile Asp Gln 65 70
50		Phe Leu Tyr Phe Asp Leu Ile Tyr Ser Ile 75 80
35		Lys Asp Thr Lys Leu Gly Asp Tyr Asp Asn 85 90
		Val Arg Val Glu Phe Lys Asn Lys Asp Leu 95 100
40		Ala Asp Lys Tyr Lys Asp Lys Tyr Val Asp 105 110
15		Val Phe Gly Ala Asn Tyr Tyr Tyr Gln Cys 115 120
45		Tyr Phe Ser Lys Lys Thr Asn Asp Ile Asn 125 130
50		Ser His Gln Thr Asp Lys Arg Lys Thr Cys

	Met Tyr Gly Gly Val Thr Glu His Asn Gly 145 150
5	Asn Gln Leu Asp Lys Tyr Arg Ser Ile Thr 155 160
	Val Arg Val Phe Glu Asp Gly Lys Asn Leu 165 170
10	Leu Ser Phe Asp Val Gln Tyr Asn Lys Lys 175 180
15	Lys Val Thr Ala Gln Glu Leu Asp Tyr Leu 185 190
13	Thr Arg His Tyr Leu Val Lys Asn Lys Lys 195 200
20	Leu Tyr Glu Phe Asn Asn Ser Pro Tyr Glu 205 210
	Thr Gly Tyr Ile Lys Phe Ile Glu Asn Gln 215 220
25	Asn Phe Trp Tyr Asp Met Met Pro Ala Pro 225 230
30	Gly Asp Lys Phe Ala Gln Ser Lys Tyr Leu 235 240
50	Met Met Tyr Asn Asp Asn Lys Met Val Asp 245 250
35	Ser Lys Asp Val Lys Leu Glu Val Tyr Leu 255 260
	Thr Thr Lys Lys 265
40	(8) INFORMATION FOR SEQUENCE ID NO:7:
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1712 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Unknown
45	(D) TOPOLOGY: Unknown (ii) Molecule type: DNA
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
50	GAACTAGGTA GAAAAATAAT TATGAGAAAA CACTATGTTG
	TTAAAGATGT TTTCGTATAT AAGTTTAGGT GATGTATAGT 8

	TACTTAATTT TAAAAGCATA ACTTAATTAA TATAAATAAC	120
5	ATGAGATTAT TAAATATAAT TAAGTTTCTT TTAATGTTTT	160
	TTTAATTGAA TATTTAAGAT TATAACATAT ATTTAAAGTG	200
	TATCTAGATA CTTTTTGGGA ATGTTGGATA AAGGAGATAA	240
10	AAAATGTATA AGAGATTATT TATTTCACAT GTAATTTTGA	280
	TATTCGCACT GATATTAGTT ATTTCTACAC CCAACGTTTT	320
15	AGCAGAGAGT CAACCAGATC CTAAACCAGA TGAGTTGCAC	360
.5	AAATCGAGTA AATTCACTGG TTTGATGGAA AATATGAAAG	400
	TTTTGTATGA TGATAATCAT GTATCAGCAA TAAACGTTAA	440
20	ATCTATAGAT CAATTTCGAT ACTTTGACTT AATATATTCT	480
	ATTAAGGACA CTAAGTTAGG GAATTATGAT AATGTTCGAG	520
25	TCGAATTTAA AAACAAAGAT TTAGCTGATA AATACAAAGA	560
	TAAATACGTA GATGTGTTTG GAGCTAATGC TTATTATCAA	600
	TGTGCTTTTT CTAAAAAAAC GAATGATATT AATTCGCATC	640
30	AAACTGACAA ACGAAAAACT TGTATGTATG GTGGTGTAAC	680
	TGAGCATAAT GGAAACCAAT TAGATAAATA TAGAAGTATT	720
35	ACTGTTCGGG TATTTGAAGA TGGTAAAAAT TTATTATCTT	760
	TTGACGTACA AACTAATAAG AAAAAGGTGA CTGCTCAAGA	800
	ATTAGATTAC CTAACTCGTC ACTATTTGGT GAAAAATAAA	840
40	AAACTCTATG AATTTAACAA CTCGCCTTAT GAAACGGGAT	880
	ATATTAAATT TATAGAAAAT GAGAATAGCT TTTGGTATGA	920
45	CATGATGCCT GCACCAGGAG ATAAATTTGA CCAATCTAAA	960
	TATTTAATGA TGTACAATGA CAATAAAATG GTTGATTCTA	1000
	AAGATGTGAA GATTGAAGTT TATCTTACGA CAAAGAAAAA	1040
50	GTGAAATTAT ATTTTAGAAA AGTAAATATG AAGAGTTAGT	1080
	AATTAAGGCA GGCACTTATA GAGTACCTGC CTTTTCTAAT	1120

	ATTATTTAGT TATAGTTATT TTTGTTATAT CTCTCTGATT	1160
5	TAGCATTAAC CCCTTGTTGC CATTATAGTT TTCACCAACT	1200
	TTAGCTGAAA TTGGGGGATC ATTTTTATCT TTACTATGGA	1240
	TAGTTACTGT GTCGCCGTTT TTAACGATTT GTTTCTCTTT	1280
10	TAATTIGICA GITAATTITI TCCATGCATC ATTIGCGTCA	1320
	AACCTATTTC CATTTGGATT TATTCTTGAC AAATCAATTC	1360
15	TTTTAACACT ATCGGTATTA ATCGGCTTGT TATTAAAATT	1400
	ACTAAGTTCA TCTAAATCAG CTGTACCCGT AATACTACTT	1440
	TCGCCACCAT TATTTAAATT GTACGTAACA CCAACTGTCT	1480
20	CATTTGCTGT TTTATCGATA ATATTTGCTT CTTTCAAAGC	1520
	ATCTCTTACA TTTTTCCATA AGTCTCTATC TGTTATTTCA	1560
25	GAAGCCTTTG CAACGTTATT AATACCATTA TAATTTGAAG	1600
	AAGAATGAAA ACCTGAACCT ACTGTTGTTA AAACTAAAGC	1640
	ACTTGCTATC AATGTTCTTG TTAATAGTTT TTTATTCATT	1680
30	TTATTTCTC CTATAACTTA TTTGCAATCG AT	1712
	(9) INFORMATION FOR SEQUENCE ID NO:8:	
35	(i) SEQUENCE CHARACTERISTICS:	
33	(A) LENGTH: 265 (B) TYPE: Amino Acid	
	(C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown	
10	<pre>(ii) Molecule type: Peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO</pre>	٠
	Met Tyr Lys Arg Leu Phe Ile Ser His Val	
	5 10	
15	Ile Leu Ile Phe Ala Leu Ile Leu Val Ile 15 20	
	Ser Thr Pro Asn Val Leu Ala Glu Ser Gln	
0	25 30	
	Pro Asp Pro Lys Pro Asp Glu Leu His Lys	

					35					40
5	Se	er Se	r Lys	Phe	Thr 45	Gly	Leu	Met	Glu	Asn 50
,	· Me	t Ly	s Val	. Leu	Tyr 55	Asp	Asp	Asn	His	Va]
10	Se	er Ala	a Ile	: Asn	Val 65	Lys	Ser	Ile	Asp	Glr 70
	Ph	e Ar	g Tyr	Phe	Asp 75	Leu	Ile	Туг	Ser	Ile 80
15	Ly	s Ası	o Thr	Lys	Leu 85	Gly	Asp	Tyr	Asp	Asn 90
20	Va	l Arg	y Val	Glu	Phe 95	Lys	Asn	Lys	Asp	Leu 100
20	Al	a Ası) Lys	Tyr	Lys 105	Asp	Lys	Tyr	Val	Asp 110
25	Va	l Phe	e Gly	Ala	Asn 115	Ala	Tyr	Tyr	Gln	Cys 120
	Al	a Phe	e Ser	Lys	Lys 125	Thr	Asn	Asp	Ile	Asn 130
30	Se	r His	Gln	Thr	Asp 135	Lys	Arg	Lys	Thr	Cys 140
35	Me	t Tyr	Gly	Gly	Val 145	Thr	Glu	His	Asn	Gly 150
	Ası	n Gln	Leu	Asp	Lys 155	Tyr	Arg	Ser	Ile	Thr 160
40	Va.	l Arg	Val	Phe	Glu 165	Asp	Gly	Lys	Asn	Leu 170
	Let	ı Ser	Phe	Asp	Val 175	Gln	Tyr	Asn	Lys	Lys 180
45	Lys	Val	Thr	Ala	Gln 185	Glu	Leu	Asp	Tyr	Leu 190
50	Thi	Arg	His	Tyr	Leu 195	Val	Lys	Asn	Lys	Lys 200
30	Lev	ı Tyr	Glu	Phe	Asn 205	Asn	Ser	Pro	Tyr	Glu 210

	Thr Gly Tyr Ile Lys Phe Ile Glu Asn Gln 215 220
5	Asn Phe Trp Tyr Asp Met Met Pro Ala Pro 225 230
10	Gly Asp Lys Phe Asp Gln Ser Lys Tyr Leu 235 240
10	Met Met Tyr Asn Asp Asn Lys Met Val Asp 245 250
15	Ser Lys Asp Val Lys Leu Glu Val Tyr Leu 255 260
	Thr Thr Lys Lys Lys 265
20	(10) INFORMATION FOR SEQUENCE ID NO:9:
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1388 (B) TYPE: Nucleic Acid
25	(C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: DNA
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
30	ATGAGTCAAC CAGATCCTAA ACCAGATGAG TTGCACAAAT 4
	CGAGTAAATT CACTGGTTTG ATGGAAAATA TGAAAGTTTT 8
35	GTATGATGAT AATCATGTAT CAGCAATAAA CGTTAAATCT 12
33	ATAGATCAAT TTCGATACTT TGACTTAATA TATTCTATTA 16
	AGGACACTAA GTTAGGGAAT TATGATAATG TTCGAGTCGA 20
40	ATTTAAAAAC AAAGATTTAG CTGATAAATA CAAAGATAAA 240
	TACGTAGATG TGTTTGGAGC TAATGCTTAT TATCAATGTG 280
45	CTTTTTCTAA AAAAACGAAT GATATTAATT CGCATCAAAC 320
	TGACAAACGA AAAACTTGTA TGTATGGTGG TGTAACTGAG 360
	CATAATGGAA ACCAATTAGA TAAATATAGA AGTATTACTG 400
50	TTCGGGTATT TGAAGATGGT AAAAATTTAT TATCTTTTGA 441

	CGTACAAACT AATAAGAAAA AGGTGACTGC TCAAGAATTA	480
	GATTACCTAA CTCGTCACTA TTTGGTGAAA AATAAAAAAC	520
5	TCTATGAATT TAACAACTCG CCTTATGAAA CGGGATATAT	560
	TAAATTTATA GAAAATGAGA ATAGCTTTTG GTATGACATG	600
10	ATGCCTGCAC CAGGAGATAA ATTTGACCAA TCTAAATATT	640
10	TAATGATGTA CAATGACAAT AAAATGGTTG ATTCTAAAGA	680
	TGTGAAGATT GAAGTTTATC TTACGACAAA GAAAAAGTGA	720
15	AATTATATTT TAGAAAAGTA AATATGAAGA GTTAGTAATT	760
	AAGGCAGGCA CTTATAGAGT ACCTGCCTTT TCTAATATTA	800
20	TTTAGTTATA GTTATTTTTG TTATATCTCT CTGATTTAGC	840
	ATTAACCCCT TGTTGCCATT ATAGTTTTCAC CAACTTTAG	880
	CTGAAATTGG GGGATCATTT TTATCTTTAC TATGGATAGT	920
25	TACTGTGTCG CCGTTTTTAA CGATTTGTTT CTCTTTTAAT	960
	TTGTCAGTTA ATTTTTTCCA TGCATCATTT GCGTCAAACC	1000
30	TATTTCCATT TGGATTTATT CTTGACAAAT CAATTCTTTT	1040
	AACACTATCG GTATTAATCG GCTTGTTATT AAAATTACTA	1080
	AGTTCATCTA AATCAGCTGT ACCCGTAATA CTACTTTCGC	1120
35	CACCATTATT TAAATTGTAC GTAACACCAA CTGTCTCATT	1160
	TGCTGTTTTA TCGATAATAT TTGCTTCTTT CAAAGCATCT	1200
40	CTTACATTIT TCCATAAGTC TCTATCTGTT ATTTCAGAAG	1240
•	CCTTTGCAAC GTTATTAATA CCATTATAAT TTGAAGAAGA	1280
	ATGAAAACCT GAACCTACTG TTGTTAAAAC TAAAGCACTT	1320
45	GCTATCAATG TTCTTGTTAA TAGTTTTTTA TTCATTTTAT	1360
	TTTCTCCTAT AACTTATTTG CAATCGAT	1388
	(11) INFORMATION FOR SEQUENCE ID NO:10:	
50	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 239	

	(B) TYPE: Amino Acid (C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: Peptide	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
	Met Ser Gln Pro Asp Pro Lys Pro Asp Glu 5 10	
10	Leu His Lys Ser Ser Lys Phe Thr Gly Lys 15 20	
15	Met Glu Asn Met Lys Val Leu Tyr Asp Asp 25 30	
	Asn His Val Ser Ala Ile Asn Val Lys Ser 35 40	
20	Ile Asp Gln Phe Arg Tyr Phe Asp Leu Ile 45 50	
	Tyr Ser Ile Lys Asp Thr Lys Leu Gly Asn 55 60	
25	Tyr Asp Asn Val Arg Val Glu Phe Lys Asn 65 70	
30	Lys Asp Leu Ala Asp Lys Tyr Lys Asp Lys 75 80	
	Tyr Val Asp Val Phe Gly Ala Asn Ala Tyr 85 90	
35	Tyr Gln Cys Ala Phe Ser Lys Lys Thr Asn 95 100	
	Asp Ile Asn Ser His Gln Thr Asp Lys Arg 105 110	
40	Lys Thr Cys Met Tyr Gly Gly Val Thr Glu 115 120	
45	His Asn Gly Asn Gln Leu Asp Lys Tyr Arg 125 130	
	Ser Ile Thr Val Arg Val Phe Glu Asp Gly 135 140	
50	Lys Asn Leu Leu Ser Phe Asp Val Gln Thr 145 150	

	Asn Lys Lys Lys Val Thr Ala Gln Glu Leu 155 160
5	Asp Tyr Leu Thr Arg His Tyr Leu Val Lys 165 170
	Asp Lys Lys Leu Tyr Glu Phe Asn Asn Ser 175 180
10	Pro Tyr Glu Thr Gly Tyr Ile Lys Phe Ile . 185 190
15	Glu Asn Glu Asn Ser Phe Trp Tyr Asp Met 195 200
15	Met Pro Ala Pro Gly Asp Lys Phe Asp Gln 205 210
20	Ser Lys Tyr Leu Met Met Tyr Asn Asp Asn 215 220
	Lys Met Val Asp Ser Lys Asp Val Lys Ile 225 230
25	Glu Val Tyr Leu Thr Thr Lys Lys Lys 235
	(12) INFORMATION FOR GROUPING TO NO. 11
30	(12) INFORMATION FOR SEQUENCE ID NO:11: (i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 731 (B) TYPE: Nucleic Acid
	(C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown
35	(ii) Molecule type: DNA
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
	TAAGGAGAAT TAAAAATGAA TAAAAAATTA CTAATGAATT 40
40	TTTTTATCGT AAGCCCTTTG TTGCTTGCGA CAACTGCTAC 80
	AGATTTTACC CCTGTTCCCT TATCATCTAA TCAAATAATC 120
45	AAAACTGCAA AAGCATCTAC AAACGATAAT ATAAAGGATT 160
	TGCTAGACTG GTATAGTAGT GGGTCTGACA CTTTTACAAA 200
	TAGTGAAGTT TTAGATAATT CCAGAGGATC TATGCGTATA 240
50	AAAAACACAG ATGGCAGCAT CAGCTTGATA ATTTTTCCGA 280

		GTCC	TATT	ГА ТА	GCCC1	CCT	TTTAC	AAAAC	GGG.	аааа	GT	320
		TGAC	TTAA	AC AC	AAAA	GAA	СТААА	AAAAG	CCA	ACATA	.CT	360
5		AGCG	AAGG	AA CI	TATAT	CCA	TTTCC	AAATA	AGT	GCCT	TA	400
		CAAA	TACT	GA AA	AATTA	CCT .	ACTCC	AATAC	AAC	TACCT	TT	440
10		AAAA	GTTA	AG GT	TCATO	GTA .	AAGAT	AGCCC	CTT	AAAGT	'AT	480
		GGGC	CAAA	ST TO	GATA?	AAA.	ACAAT	TAGCI	'ATA	rcaac	TT	520
		TAGA	CTTT	GA AA	TTCGI	CAT	CAGCT	AACTO	: AAA	FACAT	GG	560
15		ATTA	TATC	T TC	AAGCG	ATA	AAACG	GGTGG	TTA'	r t gga	AA	600
		ATAA	ÇAAT	GA AT	GACGG	SATC (CACAT	ATCAA	AGT	GATTT	AT	640
20		CTAA	AAAG'	rt tg	AATAC	AAT I	ACTGA	AAAAC	CAC	CTATA	AA	680
		TATT	GATG	AA AT	AAAAA	CTA '	TAGAA	GCAGA	AAT	TTAAT	AA .	720
			CCAC									731
30	(13)	INFOR		SEQU (A) (B) (C)	ENCE LEN TYP STR	CHA GTH: E: A ANDE	RACT 233 mino DNES	ERIS	TICS d nkno	:		
<i>5</i> 0		(ii) 1		cule	typ	e: P	epti	de		EQ I	D NO	:12:
35		Met	Asn	Lys	Lys	Leu 5	Leu	Met	Asn	Phe	Phe 10	
		Ile	Val	Ser	Pro	Leu 15	Leu	Leu	Ala	Thr	Thr 20	
40		Ala	Thr	Asp	Phe	Thr 25	Pro	Val	Pro	Leu	Ser 30	
45		Ser	Asn	Gln	Ile	Ile 35	Lys	Thr	Ala	Lys	Ala 40	
_		Ser	Thr	Asn	Asp	Asn 45	Ile	Lys	Asp	Leu	Leu 50	
50		Asp	Trp	Tyr	Ser	Ser 55	Gly	Ser	Asp	Thr	Phe 60	

	,	Thr	Asn	Ser	Glu	Val 65	Leu	Asp	Asn	Ser	7 Arg
5		Gly	Ser	Met	Arg	Ile 75	Lys	Asn	Thr	Asp	Gl ₃
	\$	Ser	Ile	Ser	Lys	Ile 85	Ile	Phe	Pro	Ser	Pro 90
10	1	ſyr	Tyr	Ser	Pro	Ala 95	Phe	Thr	Lys	Gly	G1: 100
15	I	љуs	Val	Asp	Leu	Asn 105	Thr	Lys	Arg	Thr	Lys 110
••	I	уѕ	Ser	Gln	His	Thr 115	Ser	Gly	Thr	Tyr	Ile 120
20	F	lis	Phe	Gln	Ile	Ser 125	Gly	Val	Thr	Asn	Th: 130
	G	lu	Lys	Leu	Pro	Thr 135	Pro	Ile	Glu	Leu	Pro 140
25	I	eu	Lys	Val	Lys	Val 145	His	Gly	Lys	Asp	Ser 150
30	F	ro	Leu	Lys	Tyr	Gly 155	Pro	Lys	Phe	Asp	Lys 160
	L	ys	Gln	Leu	Ala	Ile 165	Ser	Thr	Leu	Asp	Phe 170
35	G	lu	Ile	Arg	His	Gln 175	Leu	Thr	Gln	Ile	His 180
	G	ly	Leu	Tyr	Arg	Ser 185	Ser	Asp	Lys	Thr	Gly 190
40	G	ly	Tyr	Trp	Lys	Ile 195	Thr	Met	Asn	Asp	Gly 200
45	s	er	Thr	Tyr	Gln	Ser 205	Asp	Leu	Ser	Lys	Lys 210
73	P	he	Glu	Туr	Asn	Thr 215	Glu	Lys	Pro	Pro	Ile 220
50	A	sn	Ile	Asp	Glu	Ile 225	Lys	Thr	Ile	Glu	Ala 230
	G	lu	Ile	Asn							

	(14) INFORMATION FOR SEQUENCE ID NO:13:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1095 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: DNA	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID N	0:13:
	ATCATTAAAT ATAATTAATT TTCTTTTAAT ATTTTTTTAA	40
	TTGAATATTT AAGATTATAA GATATATTTA AAGTGTATCT	80
15	AGATACTTTT TGGGAATGTT GGATGAAGGA GATAAAAATG	120
	AATAAGAGTC GATTTATTTC ATGCGTAATT TTGATATTCG	160
20	CACTTATACT AGTTCTTTTT ACACCCAACG TATTAGCAGA .	200
	GAGCCAACCA GACCCTACGC CAGATGAGTT GCACAAAGCG	240
25	AGTAAATTCA CTGGTTTGAT GGAAAATATG AAAGTTTTAT	280
	ATGATGATCA TTATGTATCA GCAACTAAAG TTAAGTCTGT	320
	AGATAAATTT AGGGCACATG ATTTAATTTA TAACATTAGT	360
30	GATAAAAAC TGAAAAATTA TGACAAAGTG AAAACAGAGT	400
	TATTAAATGA AGGTTTAGCA AAGAAGTACA AAGATGAAGT	440
	AGTTGATGTG TATGGATCAA ATTACTATGT AAACTGCTAT	480
35	TTTTCATCCA AAGATAATGT AGGTAAAGTT ACAGGTGGCA	520
	AAACTTGTAT GTATGGAGGA ATAACAAAAC ATGAAGGAAA	560
40	CCACTTTGAT AATGGGAACT TACAAAATGT ACTTATAAGA	600
	GTTTATGAAA ATAAAAGAAA CACAATTTCT TTTGAAGTGC	640
45	AAACTGATAA GAAAAGTGTA ACAGCTCAAG AACTAGACAT	680
,,	AAAAGCTAGG AATTTTTTAA TTAATAAAAA AAATTTGTAT	720
	GAGTTTAACA GTTCACCATA TGAAACAGGA TATATAAAAT	760
50	TTATTGAAAA TAACGGCAAT ACTTTTTGGT ATGATATGAT	800

		GCCTGCACCA GGCGATAAGT TTGACCAATC TAAATATTTA 840
		ATGATGTACA ACGACAATAA AACGGTTGAT TCTAAAAGTG 880
5		TGAAGATAGA AGTCCACCTT ACAACAAAGA ATGGATAATG 920
		TTAATCCGAT TTTGATATAA AAAGTGAAAG TATTAGATAT 960
10		ATTTGAAAGG TAAGTACTTC GGTGCTTGCC TTTTTAGGAT 1000
10		GCATATATAT AGATTAAACC GCACTTCTAT ATTAATAGAA 1040
		AGTGCGGTTA TTTATACACT CAATCTAAAC TATAATAATT 1080
15		GGAATCATCT TCAAA 1095
	(15)	INFORMATION FOR SEQUENCE ID NO:14:
20		(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 266 (B) TYPE: Amino Acid (C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: Peptide
25		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
		Met Asn Lys Ser Arg Phe Ile Ser Cys Val 5 10
30		Ile Leu Ile Phe Ala Leu Ile Leu Val Leu 15 20
35		Phe Thr Pro Asn Val Leu Ala Glu Ser Gln 25 30
		Pro Asp Pro Thr Pro Asp Glu Leu His Lys 35 40
40		Ala Ser Lys Phe Thr Gly Leu Met Glu Asn 45 50
		Met Lys Val Leu Tyr Asp Asp His Tyr Val 55 . 60
45		Ser Ala Thr Lys Val Lys Ser Val Asp Lys 65 70
50		Phe Arg Ala His Asp Leu Ile Tyr Asn Ile 75 80
		Ser Asp Lys Lys Leu Lys Asp Tyr Asp Lyc

					85					90
5	Val	Lys	Thr	Glu	Leu 95	Leu	Asr	Glu	Gly	Let 100
,	Ala	Lys	Lys	Тут	Lys 105	Asp	Glu	Val	Val	Asp 110
10	Val	Tyr	Gly	Ser	Asn 115	Туг	Туг	Val	Asn	Cys 120
	Tyr	Phe	Ser	Ser	Lys 125	Asp	Asn	Val	Gly	Lys 130
15	Val	Thr	Gly	Gly	Lys 135	Thr	Cys	Met	тух	Gly 140
20	Gly	Ile	Thr	Lys	His 145	Glu	Gly	Asn	His	Phe 150
20	Asp	Asn	Gly	Asn	Leu 155	Gln	Asn	Val	Leu	Ile 160
25	Arg	Val	Tyr	Glu	Asn 165	Lys	Arg	Asn	Thr	Ile 170
	Ser	Phe	Glu	Val	Gln 175	Thr	Asp	Lys	Lys	Ser 180
30	Val	Thr	Ala	Gln	Glu 185	Leu	Asp	Ile	Lys	Ala 190
35	Arg	Asn	Phe	Leu	Ile 195	Asn	Lys	Lys	Asn	Leu 200
	Tyr	Glu	Phe	Asn	Ser 205	Ser	Phe	Tyr	Glu	Thr 210
40	Gly	Tyr	Ile	Lys	Phe 215	Ile	Glu	Asn	Asn	Gly 220
	Asn	Thr	Phe	Trp	Tyr 225	Asp	Met	Met	pro	Ala 230
45	Pro	Gly	Asp	Lys	Phe 235	Asp	Gln	Ser	Lys	Tyr 240
50	Leu	Met	Met	Tyr	Asn 245	Asp	Asn	Lys	Thr	Val 250
	Asp	Ser	Lys	Ser	Val 255	Lys	Ile	Glu	Val	His 260

Leu Thr Thr Lys Asn Gly 265

5	(16)	INFORMATION FOR SEQUENCE ID NO:15:	
10	-	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1837 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (ii) Molecule type: DNA	
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15	5:
15		TCATGTTTGA CAGCTTATCA TCGATAAGCT TACTTTTCGA	40
13		ATCAGGTCTA TCCTTGAAAC AGGTGCAACA TAGATTAGGG	80
		CATGGAGATT TACCAGACAA CTATGAACGT ATATACTCAC	120
20		ATCACGCAAT CGGCAATTGA TGACATTGGA ACTAAATTCA	160
		ATCAATTTGT TACTAACAAG CAACTAGATT GACAACTAAT 2	200
25		TCTCAACAAA CGTTAATTTA ACAACATTCA AGTAACTCCC 2	40
25		ACCAGCTCCA TCAATGCTTA CCGTAAGTAA TCATAACTTA 2	80
		CTANANCCTT GTTACATCAN GGTTTTTTCT TTTTGTCTTG	20
30		TTCATGAGTT ACCATAACTT TCTATATTAT TGACAACTAA	60
		ATTGACAACT CTTCAATTAT TTTTCTGTCT ACTCAAAGTT	100
25		TTCTTCATTT GATATAGTCT AATTCCACCA TCACTTCTTC	40
35		CACTCTCTC ACCGTCACAA CTTCATCATC TCTCACTTTT 4	80
		TCGTGTGGTA ACACATAATC AAATATCTTT CCGTTTTTAC 5	20
40		GCACTATCGC TACTGTGTCA CCTAAAATAT ACCCCTTATC 5	60
		AATCGCTTCT TTAAACTCAT CTATATATAA CATATTTCAT 6	00
		CCTCCTACCT ATCTATTCGT AAAAAGATAA AAATAACTAT 6	40
45		TGTTTTTTT GTTATTTTAT AATAAAATTA TTAATATAAG 6	80
		TTAATGTTTT TTAAAAATAT ACAATTTTAT TCTATTTATA 7	20
50		GTTAGCTATT TTTTCATTGT TAGTAATATT GGTGAATTGT 7	60

	i	AATAACCTTT	TTAAATCTAG	AGGAGAACCC	AGATATAAAA	800
5		rggaggaata	TTAATGGAAA	АСААТААААА	AGTATTGAAG	840
,	i	AAAATGGTAT	TTTTTGTTT	AGTGACATTT	CTTGGACTAA	880
	·	CAATCTCGCA	AGAGGTATTT	GCTCAACAAG	ACCCCGATCC	920
10	į	AAGCCAACTT	CACAGATCTA	GTTTAGTTAA	AAACCTTCAA	960
	i	AATATATATT	TTCTTTATGA	GGGTGACCCT	GTTACTCACG	1000
15	1	AGAATGTGAA	ATCTGTTGAT	CAACTTAGAT	CTCACGATTT	1040
13	1	АТАТАТААТ	GTTTCAGGGC	CAAATTATGA	ТАААТТАААА	1080
	1	ACTGAACTTA	AGAACCAAGA	GATGGCAACT	TTATTTAAGG	1120
20	,	ATAAAAACGT	TGATATTTAT	GGTGTAGAAT	ATTACCATCT	1160
	C	TGTTATTTA	TGTGAAAATG	CAGAAAGGAG	TGCATGTATC	1200
25	1	PACGGAGGG	TAACAAATCA	TGAAGGGAAT	CATTTAGAAA	1240
23	מ	тсставава	GATAGTCGTT	AAAGTATCAA	TCGATGGTAT	1280
	C	CAAAGCCTA	TCATTTGATA	TTGAAACAAA	TAAAAAAATG	1320
30		STAACTGCTC	AAGAATTAGA	CTATAAAGTT	AGAAAATATC	1360
	7	TACAGATAA	TAAGCAACTA	TATACTAATG	GACCTTCTAA	1400
35	P	TATGAAACT	GGATATATAA	AGTTCATACC	TAAGAATAAA	1440
55	G	CAAAGTTTTT	GGTTTGATTT	TTTCCCTGAA	CCAGAATTTA	1480
	c	TCAATCTAA	ATATCTTATG	ATATATAAAG	ATAATGAAAC	1520
40	G	CTTGACTCA	AACACAAGCC	AAATTGAAGT	CTACCTAACA	1560
	А	CCAAGTAAC	TTTTTGCTTT	TGGCAACCTT	ACCTACTGCT	1600
45	G	GATTTAGAA	ATTTTATTGC	AATTCTTTTA	ттаатстааа	1640
40	А	ACCGCTCAT	TTGATGAGCG	GTTTTGTCTT	ATCTAAAGGA	1680
	G	CTTTACCTC	CTAATGCTGC	AAAATTTTAA	ATGTTGGATT	1720
50	т	TTGTATTTG	TCTATTGTAT	TTGATGGGTA	ATCCCATTTT	1760
	т	CGACAGACA	TCGTCGTGCC	ACCTCTAACA	ССААААТСАТ	1800

(17) INFORMATION FOR SEQUENCE ID NO:16: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 251 (B) TYPE: Amino Acid (C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (II) Molecule type: Peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: Met Glu Asn Asn Lys Lys Val Leu Lys Lys 5 10 Met Val Phe Phe Val Leu Val thr Phe Leu 15 20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 45 40 26 Arg Ser Ser Leu Tyr Glu Gly Asp Pro Val 55 60 The His Glu Asn Val Lys Ser Val Asp Gln 65 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 95 105 Val Glu Tyr Tyr His Leu Cys Tyr Leu Cys		AGACAGGAGC TTGTAGCTTA GCAACTATTT TATCGTC 1837
5 (i) SEQUENCE CHARACTERISTICS:		(17) INFORMATION FOR SEQUENCE ID NO:16:
(B) TYPE: Amino Acid (C) STRANDEENESS: Unknown (D) TOPOLOGY: Unknown (E) Topolog: Unknown (E) Topological Experiments (E) Topological Experiments (E)	5	
(C) STRANDEDNESS: Unknown (D) TOPOLOGY: Unknown (E) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (E) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (D) TOPOLOGY: Unknown (E) TOPOLOGY: Unknown (D) Topology: Unknown (E) Topology: Unkn		
(ii) Molecule type: Peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: Met Glu Asn Asn Lys Lys Val Leu Lys Lys 10 Met Val Phe Phe Val Leu Val thr Phe Leu 15 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 25 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 35 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 55 Thr His Glu Asn Val Lys Ser Val Asp Gln 65 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 95 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105		(C) STRANDEDNESS: Unknown
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: Met Glu Asn Asn Lys Lys Val Leu Lys Lys 10 15 Met Val Phe Phe Val Leu Val thr Phe Leu 20 20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 35 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 30 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 30 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 35 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	10	
Met Val Phe Phe Val Leu Val thr Phe Leu 20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110		
Met Val Phe Phe Val Leu Val thr Phe Leu 20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110		Met Glu Asn Asn Lys Lys Val Leu Lys Lys
Met Val Phe Phe Val Leu Val thr Phe Leu 20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asp Lys Asn Val Asp Ile Tyr Gly 105 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	15	
20 Gly Leu Thr Ile Ser Gln Glu Val Phe Ala 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asp Lys Asn Val Asp Ile Tyr Gly 110 45		
25 30 Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110 45		15 20
Gln Gln Asp Pro Asp Pro Ser Gln Leu His 40 25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 50 Ile Tyr Phe Leu Tyr Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	20	
25 Arg Ser Ser Leu Val Lys Asn Leu Gln Asn 45		
11e Tyr Phe Leu Tyr Glu Gly Asp Pro Val 55 Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 Asp Lys Leu Lys Thr 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 Phe Lys Asp Lys Asn Cln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110		35 40
11e Tyr Phe Leu Tyr Glu Gly Asp Pro Val 55 Glu Gly Asp Pro Val 60 Thr His Glu Asn Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 Asp Lys Leu Lys Thr 80 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 Phe Lys Asp Lys Asn Cln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	25	Arg Ser Ser Leu Val Lys Asn Leu Gln Asn
Thr His Glu Asn Val Lys Ser Val Asp Gln 65 Val Lys Ser Val Asp Gln 70 Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 Leu Lys Thr 85 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105		
Thr His Glu Asn Val Lys Ser Val Asp Gln Thr His Glu Asn Val Lys Ser Val Asp Gln To Leu Arg Ser His Asp Leu Ile Tyr Asn Val To Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105		
Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	30	55 60
Leu Arg Ser His Asp Leu Ile Tyr Asn Val 75 Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105		
Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 85 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110		
Ser Gly Pro Asn Tyr Asp Lys Leu Lys Thr 90 40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110	35	
40 Glu Leu Lys Asn Gln Glu Met Ala Thr Leu 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 110 45		
95 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105 110		
95 100 Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly 105 110	40	Glu Leu Lys Asn Gln Glu Met Ala Thr Leu
45		
45		Phe Lys Asp Lys Asn Val Asp Ile Tyr Gly
Val Glu Tyr Tyr His Leu Cys Tyr Leu Cys	45	105 110
		Val Glu Tyr Tyr His Leu Cys Tyr Leu Cys
115 120		
Glu Asn Ala Glu Arg Ser Ala Cys Ile Tyr 50 125 130	50	

WO 00/09154 PCT/US98/16766

	Gly	Gly	Val	Thr	Asn 135	His	Glu	Gly	Asn	His 140
5	Leu	Glu	Ile	Pro	Lys 145	Lys	Ile	Val	Val	Lys 150
	Val	Ser	Ile	Asp	Gly 155	Ile	Gln	Ser	Leu	Ser 160
10	Phe	Asp	Ile	Glu	Thr 165	Asn	Lys	Lys	Met	Val 170
15	Thr	Ala	Gln	Glu	Leu 175	Asp	Tyr	Lys	Val	Arg 180
15	Lys	Tyr	Leu	Thr	Asp 185	Asn	Lys	Gln	Leu	Туг 190
20	Thr	Asn	Gly	Pro	Ser 195	Lys	Tyr	Glu	Thr	Gly 200
	Tyr	Ile	Lys	Phe	Ile 205	Pro	Lys	Asn	Lys	Glu 210
25	Ser	Phe	Trp	Phe	Asp 215	Phe	Phe	Pro	Glu	Pro 220
30	Glu	Phe	Thr	Gln	Ser 225	Lys	Tyr	Leu	Met	Ile 230
30	Tyr	Lys	Asp	Asn	Glu 235	Thr	Leu	Asp	Ser	Asn 240
35	Thr	Ser	Gln	Ile	Glu 2 4 5	Val	Tyr	Leu	Thr	Thr 250
	Lys									

What is claimed is:

- 1. An isolated and purified superantigen toxin DNA fragment which has been altered such that binding of the encoded altered toxin to either the MHC class II or T cell antigen receptor is altered.
- An isolated and purified DNA fragment according to claim 1, wherein said superantigen toxin
 is Staphylococcal enterotoxin A having the sequence of SEQ ID NO:1 or a portion thereof, or an allelic portion thereof.
- 3. An isolated and purified DNA fragment
 15 according to claim 1, wherein said superantigen toxin
 is Staphylococcal enterotoxin A having the sequence of
 SEQ ID NO:3 or a portion thereof, or an allelic
 portion thereof.
- 4. An isolated and purified DNA fragment according to claim 1, wherein said superantigen toxin is Staphylococcal enterotoxin B having the sequence of SEQ ID NO:5 or a portion thereof, or an allelic portion thereof.
- 25

- 5. An isolated and purified DNA fragment according to claim 1, wherein said superantigen toxin is Staphylococcal enterotoxin B having the sequence of SEQ ID NO:7 or a portion thereof, or an allelic portion thereof.
- 6. An isolated and purified DNA fragment according to claim 1, wherein said superantigen toxin is Staphylococcal enterotoxin B having the sequence of

SEQ ID NO:9 or a portion thereof, or an allelic portion thereof.

- 7. An isolated and purified DNA fragment according to claim 2, wherein said fragment encodes the amino acid sequence of SEQ ID NO:2 or a portion thereof, or an allelic portion thereof.
- 8. An isolated and purified DNA fragment according to claim 3, wherein said fragment encodes the amino acid sequence of SEQ ID NO:4 or a portion thereof, or an allelic portion thereof.
- 9. An isolated and purified DNA fragment according to claim 4, wherein said fragment encodes the amino acid sequence of SEQ ID NO:6 or a portion thereof, or an allelic portion thereof.
- 10. An isolated and purified DNA fragment 20 according to claim 5, wherein said fragment encodes the amino acid sequence of SEQ ID NO:8 or a portion thereof, or an allelic portion thereof.
- 11. An isolated and purified DNA fragment according to claim 6, wherein said fragment encodes the amino acid sequence of SEQ ID NO:10 or a portion thereof, or an allelic portion thereof.
 - 12. A recombinant DNA construct comprising:

(i) a vector, and

- (ii) an isolated and purified altered superantigen toxin DNA fragment according to claim 1.
- 13. A recombinant DNA construct according to 35 claim 12, wherein said DNA fragment has the sequence

according to SEQ ID NO:1 or a portion thereof, or an allelic portion thereof.

- 14. A recombinant DNA construct according to 5 claim 12, wherein said DNA fragment has the sequence according to SEQ ID NO:3 or a portion thereof, or an allelic portion thereof.
- 15. A recombinant DNA construct according to claim 12, wherein said DNA fragment has the sequence according to SEQ ID NO:5 or a portion thereof, or an allelic portion thereof.
- 16. A recombinant DNA construct according to claim 12, wherein said DNA fragment has the sequence according to SEQ ID NO:7 or a portion thereof, or an allelic portion thereof.
- 17. A recombinant DNA construct according to 20 claim 12, wherein said DNA fragment has the sequence according to SEQ ID NO:9 or a portion thereof, or an allelic portion thereof.
- 18. The recombinant DNA construct according 25 to claim 13, wherein said DNA fragment encodes the amino acids sequence specified in SEQ ID NO:2.
- 19. The recombinant DNA construct according to claim 14, wherein said DNA fragment encodes the amino acids sequence specified in SEQ ID NO:4.
 - 20. The recombinant DNA construct according to claim 15, wherein said DNA fragment encodes the amino acids sequence specified in SEQ ID NO:6.

21. The recombinant DNA construct according to claim 16, wherein said DNA fragment encodes the amino acids sequence specified in SEQ ID NO:8.

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- 22. The recombinant DNA construct according to claim 17, wherein said DNA fragment encodes the amino acids sequence specified in SEQ ID NO:10.
- 10 23. A recombinant DNA construct according to claim 13 wherein said construct is pETA489270P.
 - 24. A recombinant DNA construct according to claim 14 wherein said construct is pETA489270C.

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- 25. A recombinant DNA construct according to claim 15 wherein said construct is pETB2360210.
- 26. A recombinant DNA construct according to claim 16 wherein said construct pETB899445P.
 - 27. A recombinant DNA construct according to claim 17 wherein said construct is pETB899445C.
- 25 28. A recombinant DNA construct according to claim 12, wherein said vector is an expression vector.
 - 29. A host cell transformed with a recombinant DNA construct according to claim 12.

- 30. A host cell transformed with a recombinant DNA construct according to claim 18.
- 31. A host cell transformed with a recombinant DNA construct according to claim 19.

- 32. A host cell transformed with a recombinant DNA construct according to claim 20.
- 5 33. A host cell transformed with a recombinant DNA construct according to claim 21.
 - 34. A host cell transformed with a recombinant DNA construct according to claim 22.
 - 35. A host cell according to claim 29, wherein said cell is prokaryotic.
- 36. A host cell according to claim 30, 15 wherein said cell is prokaryotic.
 - 37. A host cell according to claim 31, wherein said cell is prokaryotic.
- 38. A host cell according to claim 32, wherein said cell is prokaryotic.
 - 39. A host cell according to claim 33, wherein said cell is prokaryotic.
 - 40. A host cell according to claim 34, wherein said cell is prokaryotic.
- 41. A method for producing altered

 30 superantigen toxin comprising culturing the cells according to claim 29, under conditions such that said DNA fragment is expressed and said superantigen toxin is thereby produced, and isolating said superantigen toxin.

- 42. A method for producing altered superantigen toxin comprising culturing the cells adcording to claim 30, under conditions such that said DNA fragment is expressed and said superantigen toxin is, thereby produced, and isolating said superantigen toxin.
- 43. A method for producing altered

 10 superantigen toxin comprising culturing the cells according to claim 31, under conditions such that said DNA fragment is expressed and said superantigen toxin is thereby produced, and isolating said superantigen toxin.

- 44. A method for producing altered superantigen toxin comprising culturing the cells according to claim 32, under conditions such that said DNA fragment is expressed and said superantigen toxin is thereby produced, and isolating said superantigen toxin.
 - 45. A method for producing altered superantigen toxin comprising culturing the cells according to claim 33, under conditions such that said DNA fragment is expressed and said superantigen toxin is thereby produced, and isolating said superantigen toxin.
- 30 46. A method for producing altered superantigen toxin comprising culturing the cells according to claim 34, under conditions such that said DNA fragment is expressed and said superantigen toxin is thereby produced, and isolating said superantigen toxin.

- 47. An isolated and purified superantigen toxin which has been altered such that binding of the encoded altered toxin to either the MHC class II or T cell antigen receptor is altered.
 - 48. An isolated and purified superantigen toxin according to claim 47 wherein said toxin is staphylococcal enterotoxin A.

- 49. An isolated and purified superantigen toxin according to claim 47 wherein said toxin is staphylococcal enterotoxin B.
- 15 50. An altered SEA superantigen toxin peptide according to claim 48 wherein position 92 has been changed to alanine.
- 51. An altered SEA superantigen toxin
 20 peptide according to claim 48 wherein position 70 has been changed to arginine.
 - 52. An altered SEA superantigen toxin peptide according to claim 48 wherein position 48 has been changed to arginine.
 - 53. An altered SEA superantigen toxin peptide according to claim 48 wherein position 64 has been mutated to alanine.

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54. An altered SEB superantigen toxin peptide according to claim 49 wherein position 115 has been changed to alanine.

- 55. An altered SEB superantigen toxin peptide according to claim 49 wherein position 89 has been changed to alanine.
- 5 56. An altered SEB superantigen toxin peptide according to claim 49 wherein position 67 has been changed to glutamine.
- 57. An altered SEB superantigen toxin
 10 peptide according to claim 49 wherein position 94 has been changed to alanine.
- 58. An altered SEB superantigen toxin peptide according to claim 49 wherein position 61 has been changed to alanine.
 - 59. A method for the diagnosis of superantigen-associated bacterial infection comprising the steps of:
- 20 (i) contacting a sample from an individual suspected of having a superantigen-associated bacterial infection with altered superantigen toxin; and
- (ii) detecting the presence or absence of a 25 superantigen-associated bacterial infection by detecting the presence or absence of a complex formed between the altered superantigen toxin and antibodies specific therefor in the sample.
- 30 60. A method for the diagnosis of a superantigen toxin-associated bacterial infection according to claim 42 wherein the altered superantigen toxin is chosen from the group consisting of SEB, and SEA.

- 61. A superantigen toxin-associated infection diagnostic kit comprising an altered superantigen toxin according to claim 1 wherein said toxin is chosen from the group consisting of SEB, and SEA, and ancillary reagents suitable for use in detecting the presence or absence of antibodies against superantigen toxin in a mammalian sample.
- 10 62. A vaccine comprising an altered superantigen toxin according to claim 1 effective for the production of antigenic and immunogenic response resulting in the protection of a mammal against superantigen-associated bacterial infection.

63. A vaccine according to claim 62 wherein said altered superantigen toxin is chosen from the group consisting of SEB, and SEA.

- 20 64. A vaccine according to claim 63 wherein said vaccine further comprises at least one other different altered superantigen toxin chosen from the group consisting of SEB, and SEA.
- 25 65. A vaccine according to claim 62, wherein the superantigen toxin is SEB and the vaccine is identified as B899445.
- 66. A vaccine according to claim 62, wherein 30 the superantigen toxin is SEA and the vaccine is identified as A489270.
- 67. A bivalent vaccine according to claim 64 wherein said altered superantigen toxins are SEA and 35 SEB.

- 68. A bivalent vaccine according to claim 67 wherein said toxin SEA is A489270 and SEB is B899445.
- 5 69. A multivalent vaccine against superantigen-associated bacterial infections comprising a combination of altered superantigen toxins selected from the group consisting essentially of SEA, and SEB, or any portion or allelic form thereof, capable of eliciting protective antibodies against superantigen toxins in a pharmaceutically acceptable excipient in a pharmaceutically acceptable amount.
- 70. A therapeutic method for the treatment or amelioration of a superantigen-associated bacterial infection said method comprising administering to an individual in need of such treatment an effective amount of sera from individuals immunized with one of more altered superantigen toxin vaccine according to claim 62 in a pharmaceutically acceptable dose in a pharmaceutically acceptable excipient.
- 71. A therapeutic method for the treatment
 25 or amelioration of a superantigen-associated bacterial
 infection, said method comprising administering to an
 individual in need of such treatment an effective
 amount of antibodies against altered superantigen
 toxins in a pharmaceutically acceptable dose in a
 30 pharmaceutically acceptable excipient.
 - 72. A therapeutic method for the treatment or amelioration of a superantigen-associated bacterial infection, said method comprising administering to an individual in need of such treatment an effective

WO 00/09154

106

PCT/US98/16766

amount of altered superantigen toxins from staphylococcal bacteria in order to inhibit adhesion of superantigen bacterial toxin to MHC class II or T cell receptors by competitive inhibition of these interactions in a pharmaceutically acceptable dose in a pharmaceutically acceptable excipient.

73. A therapeutic method for the treatment of diseases that may not be associated directly with superantigen toxins by causing specific nonresponsiveness of T cell subsets or by expanding or stimulating specific T cell subsets, in vivo or ex vivo by use of altered superantigen toxin.

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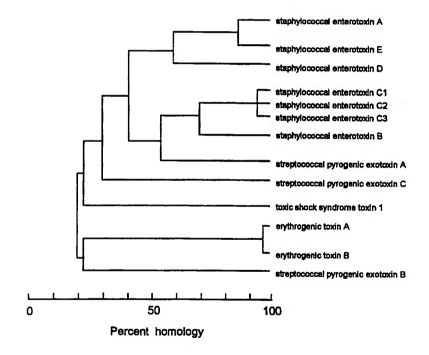
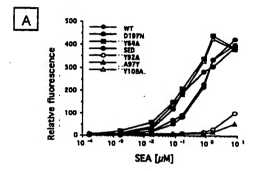
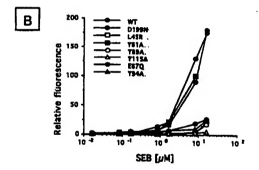


FIGURE 1





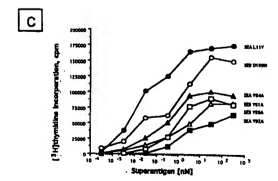
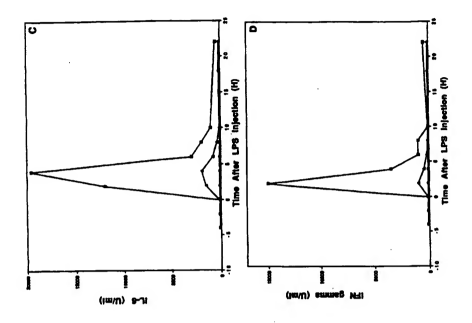
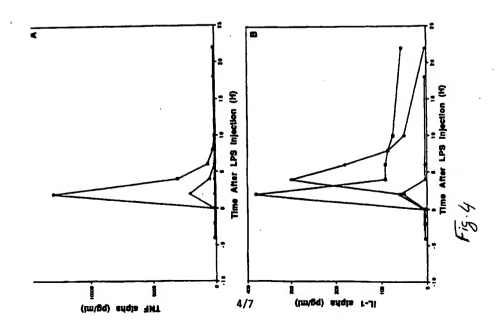


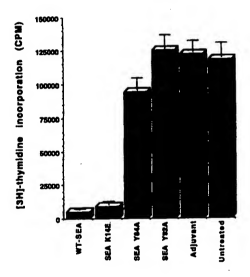
Fig. 2

SHDQP OHTILPKGPFTDHSWYNDLLV PDSKDIVDKYK. GKKVDLYGAY GYQCA GGTPNKTACH GGVTLHDNNRLTBEKK TGDQP GNTLLPKGPFTDHSWYNDLLV FENSKENAQHPK. SKNVDLYGAY GYQCA GGTDNKTACH GGVTLHDNNRLTBEKK SDDQP ENTLLPKGPFTGHPWYNDLLV LGSKDATNKYK. GKKVDLYGAY GYQCA GGTDNKTACH GGVTLHDNNRLTBEKK SDDQP ENTLLPKGPFTGHPWYNDLLV LGSKDATNKYK. GKKVDLYGAY GYQCA GGTPNKTACH GGVTLHDNNRLTBEKK SDDQP ANDLLYSIKDTGAYDNVRV FRINGALANYKK. DKVVDVFGAN YYQCYPSKRYNDINSHQTDKRKY. CH GGYTKHBGANFTRGNL SVDKF AHDLIYNI SDKKLRNYDKVKY ILLNEGLAKKYK. DEVVDVYGSN YYNCYPSSKDNVGKVTGG KT. CH GGITKHBGANFTRGNL SVDKF AHDLIYNI SDKKLRNYDKVKY ILLNEGLAKKYK. DEVVDVYGSN YYNCYPSSKDNVGKVTGG KT. CH GGITKHBGANFTRGNL SVDKF AHDLIYNI SDKKLRNYDKVKT ILLNEGLAKKYK. DEVVDVYGSN YYNCYPSSKDNVGKVTGG KT. CH GGITKHBGANFTRGNL SVDKK AHDLIYNI SDKKLRNYDKVKT ILLNEGLAKKYK. DKNVDIYGVB YHLCYLCRNAE RSACI GGYTNHBGANFTRDY SVDQL SHILLIYNSG PNYDKLKYT ILLNGGRAATLFK. DKNVDLYTKR WKSQHTSBG TYIHP.QLSGVTNH EKLPPT P
92 LYGAY G LYGAY G LYGAN Y VYGSN Y VYGSN Y VYGSN Y LYGYG YI LYGVE YI
SHDOF CHTILFKGFFTDHSWYNDLLY PDSKDIVDKYK, GKKVDLYGAY GYQCA. SHDOF CHTILFKGFFTDHSWYNDLLY PINSKEMAQHFK, SKNVDLYGAY GYQCA. STGOP KENTLLFKGFFTCHFWYNDLLY LGSKDATWKYK, GKKVDLYGAY GYQCA. STDOF YFDLIYSIKDTKLGAYDNVRY PKNKDLADKYK, GKKVDLYGAY GYQCA. SYDKF AHDLIYNISDKGLKNYDKVKY ILLNEGLAKKYK, DEVVDVYGSN YVNCYPSSKDRVGI. SVDKF AHDLIYNISDKGLKNYDKVKY ILLNEDLAKKYK, DEVVDVYGSN YVNCYPSSKDRVGI. SVDKF AHDLIYNISDKGLKNYDKVKT ILLNEDLAKKYK, DEVVDVYGSN YVNCYPSSKDRVGI. SVDKF AHDLIYNISDKGLKNYDKVKT ILLNEDLAKKYK, DEVVDVYGSN YVNCYPSSKDRVGI. SVDKF AHDLIYNISDKGLKNYDKVKT ILLNEDLAKKYK, DEVVDVYGSN YVNCYPSSKDRVGI. SVDK AHDLIXNISDKGLKNYDKYT ILLNEDLAKKYK, DKNVDIYGVE YHLCYLCRNAE. SVDK AHDLIXNISDKGLKNYDKYT ILLNEDLAKKYK, DKNVDIYGVE YHLCYLCRNAE.
TO THE TENT
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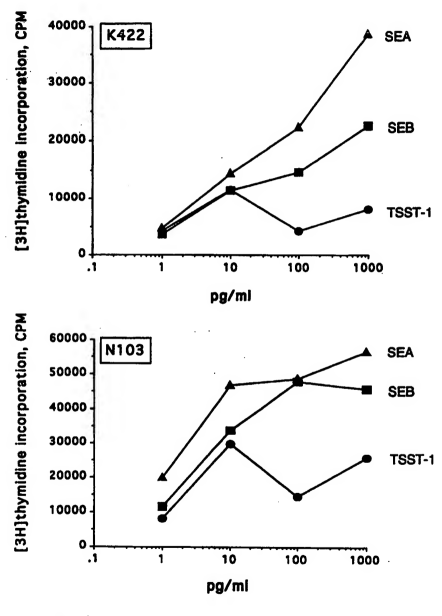
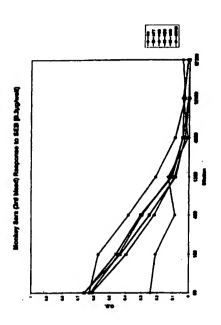
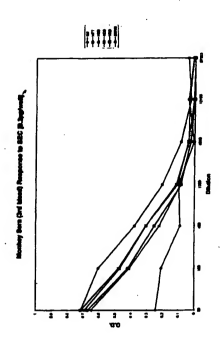
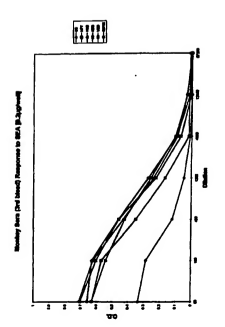


Fig. 6









7/7

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16766

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(6) :Please See Extra Sheet.						
US CL :Please See Extra Sheet. According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
Minimum documentation searched (classification system followed by classification symbols)						
U.S.: 424/184.1, 234.1, 236.1, 237.1, 243.1; 435/320.1, 252.3, 172.3, 69.1, 69.3, 70.1, 71.1, 7.1; 530/350+; 536/23.1; 436/500						
Documenta	tion searched other than minimum documentation to	the extent that such documents are included	in the fields searched			
Electronic	data base consulted during the international search	(name of data base and, where practicable	, search terms used)			
Picase Se	ee Extra Sheet.					
C. DOC	C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.			
A	DANNECKER et al. Activation of human T cells by the superantigen Staphylococcus enterotoxin B: Analysis on a cellular level. Immunobiology. 1994, Vol. 190, pages 116-126, see entire document.					
A	GONZALO et al. Expansion and clonal deletion of peripheral T cells induced by bacterial superantigen is independent of the interleukin-2 pathway. Eur. J. Immunol. 1992, Vol. 22, pages 1007-1011, see entire document.					
Y	1-73					
X Further documents are listed in the continuation of Box C. See patent family annex.						
Special categories of cited documents: A* document defining the general state of the ert which is not considered to be of particular relevance to be of particular relevance						
B° earli	earlier document published on or after the international filing data "X" document of perticular relevance; the claimed invention cannot be considered nowed or cannot be considered to involve an inventive stee.					
L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other						
special reason (as specified) *Y* document of perticular relevance; the chained investion cannot be considered to involve an investive step when the document is combined with one or more other such documents, such combination means						
P° docu						
	ate of the actual completion of the international search Date of mailing of the international search report					
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Washington, acsimile No.		Telephone No. (703) 308-0196				

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16766

nion). DOCUMENTS CONSIDERED TO BE RELEVANT	
	·
Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
BAVARI et al. Staphylococcal Enterotoxin A and toxic shock syndrome toxin compete with CD4 for human major histocompatibility complex class II binding. Infection and Immunity. February 1995, Vol. 63, No. 2, pages 423-429, see entire document.	1-73
MAHANA et al. A natural mutation of the amino acid residue at position 60 destroys staphylococcal enterotoxin A murine T-cell mitogenicity. Infection and Immunity. August 1995, Vol. 63, No. 8, pages 2826-2832, see entire document.	1-73
ULRICH et al. Bacterial superantigens in human disease: structure, function and diversity. Trends in Microbiology. 1995, Vol. 3, No. 12, pages 463-468, see entire document.	1-73
WOODY et al. Differential immune responses to Staphylococcal enterotoxin B mutations in a hydrophobic loop dominating the interface with major histocompatibility complex class II receptors. J. Infectious Diseases. 1998, Vol. 177, pages 1013-1022, see entire document.	1-73 62-69
·	
	BAVARI et al. Staphylococcal Enterotoxin A and toxic shock syndrome toxin compete with CD4 for human major histocompatibility complex class II binding. Infection and Immunity. February 1995, Vol. 63, No. 2, pages 423-429, see entire document. MAHANA et al. A natural mutation of the amino acid residue at position 60 destroys staphylococcal enterotoxin A murine T-cell mitogenicity. Infection and Immunity. August 1995, Vol. 63, No. 8, pages 2826-2832, see entire document. ULRICH et al. Bacterial superantigens in human disease: structure, function and diversity. Trends in Microbiology. 1995, Vol. 3, No. 12, pages 463-468, see entire document. WOODY et al. Differential immune responses to Staphylococcal enterotoxin B mutations in a hydrophobic loop dominating the interface with major histocompatibility complex class II receptors. J. Infectious Diseases. 1998, Vol. 177, pages 1013-1022, see

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16766

A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6):

A61K 39/00, 39/44, 39/02, 39/085; C12N 15/63+, 1/20, 1/21, 15/00; C12P 21/00+; G01N 33/53; C07K 14/00+

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

424/184.1, 234.1, 236.1, 237.1, 243.1; 435/320.1, 252.3, 172.3, 69.1, 69.3, 70.1, 71.1, 7.1; 530/350+, 536/23.1; 436/500

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, MEDLINE, BIOSIS, JAPIO, WPIDS, CABA. AGRICOLA, BIOTECHDS, CAPLUS, LIFESCI, CONFSCI, DISSABS, EMBASE

search terms: inveator names, enterotoxin A, enterotoxin B, bacterial superantigen, vaccine, dna, tsst, diagnosis, treatment, sea, seb, bivalent vaccine, staphylococcus, t-cell anergy, infection, mutant, fusion